

Rostrupomyces, a new genus to accommodate *Xerocomus sisongkhramensis*, and a new *Hemileccinum* species (Xerocomoideae, Boletaceae) from Thailand

Santhiti Vadthanarat^{1,2} , Bhavesh Raghoonundon¹ , Saisamorn Lumyong^{3,4,5}, Olivier Raspé^{1,6,7} 

¹ School of Science, Mae Fah Luang University, Chiang Rai, 57100, Thailand

² Department of Biological Science, Faculty of Science, Ubon Ratchathani University, Ubon Ratchathani, 34190, Thailand

³ Department of Biology, Faculty of Science, Chiang Mai University, Chiang Mai, 50200, Thailand

⁴ Research Center of Microbial Diversity and Sustainable Utilization, Faculty of Science, Chiang Mai University, Chiang Mai, 50200, Thailand

⁵ Academy of Science, The Royal Society of Thailand, Bangkok, Thailand

⁶ Meise Botanic Garden, Nieuwelaan 38, 1860 Meise, Belgium

⁷ Service Général de l'Enseignement Supérieur et de la Recherche Scientifique, Fédération Wallonie-Bruxelles, Brussels, Belgium

Corresponding author: Olivier Raspé (olivier.ras@mfu.ac.th)



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Abstract

A new genus, *Rostrupomyces* is established to accommodate *Xerocomus sisongkhramensis* based on multiple protein-coding genes (*atp6*, *cox3*, *tef1*, and *rpb2*) analyses of a wide taxon sampling of Boletaceae. In our phylogeny, the new genus was sister to *Rubinosporus* in subfamily Xerocomoideae, phylogenetically distant from *Xerocomus*, which was highly supported as sister to *Phylloporus* in the same subfamily Xerocomoideae. *Rostrupomyces* is different from other genera in Boletaceae by the following combination of characters: rugulose to subrugulose pileus surface, white pores when young becoming pale yellow in age, subscabrous stipe surface scattered with granulose squamules, white basal mycelium, unchanging color in any parts, yellowish brown spore print, and broadly ellipsoid to ellipsoid, smooth basidiospores. In addition, *Hemileccinum inferius*, also from subfamily Xerocomoideae, is newly described. Detailed descriptions and illustrations of the new genus and new species are presented.

Key words: *atp6*, Boletales, *cox3*, fungal diversity, multigene phylogeny, one new species, taxonomy, Tropical Asia

Introduction

Xerocomoideae Singer, which is one of the six subfamilies in Boletaceae Chevall, was established in 1945 with *Xerocomus* Quél. as the typus. At present, the subfamily consists of 12 genera, namely *Alessioporus* Gelardi, Vizzini & Simonini, *Amylotrama* Bloomfield, Davoodian, Trappe & T. Lebel, *Aureoboletus* Pouzar, *Boletellus* Murrill, *Heimioporus* E. Horak, *Hemileccinum* Šutara, *Hourangia* Xue T. Zhu & Zhu L. Yang, *Phylloporus* Quél., *Pulchroboletus* Gelardi, Vizzini & Simonini, *Rubinosporus* Vadthanarat, Raspé & Lumyong, *Veloboletus* Fechner &

Halling, and *Xerocomus* (Šutara 2008; Gelardi et al. 2014; Wu et al. 2014; Zhu et al. 2015; Wu et al. 2016; Crous et al. 2020; Lebel et al. 2022; Vadthanarat et al. 2022). The typical characters of species in this subfamily are boletoid or phylloporoid, rarely sequestrate basidiomata; dry or viscid pileus with smooth or subtomentose to tomentose pellis; absence or rarely presence of a veil; off-white, yellowish white, yellowish to yellow context; at least some basidiome parts often bluing, sometimes reddening or unchanging; smooth or ornamented stipe surface; hymenophore yellowish to yellow to bright yellow or cream to dull yellow to yellow to gray in sequestrate forms; basidiospores with bacillate, reticulate, tiny warts, pinholes, longitudinally striate, pitted ornamentations, or occasionally smooth; spore deposit with more or less olive-brown tint, rarely dark ruby (e. g. Gelardi et al. 2014; Wu et al. 2014; Zhu et al. 2015; Wu et al. 2016; Crous et al. 2020; Lebel et al. 2022; Vadthanarat et al. 2022).

Hemileccinum, one of the genera belonging to the Xerocomoideae, was established in 2008 to accommodate two *Boletus* species, namely *B. depilatus* Redeuilh and *B. impositus* Fr. In 2012, a new genus named *Corneroboletus* N.K. Zeng & Zhu L. Yang was established to accommodate *Boletus indecorus* Massee (Zeng et al. 2012). However, *Corneroboletus* was later synonymized with *Hemileccinum* (Wu et al. 2016). *Hemileccinum* currently comprises 13 species worldwide, namely *H. albidum* Mei Xiang Li, Zhu L. Yang & G. Wu, *H. brevisporum* Mei Xiang Li, Zhu L. Yang & G. Wu, *H. brunneotomentosum* (B. Ortiz) Nitson & J.L. Frank, *H. depilatum* (Redeuilh) Šutara, *H. ferrugineipes* Mei Xiang Li, Zhu L. Yang & G. Wu, *H. floridanum* J.A. Bolin, A.E. Bessette, A.R. Bessette, L.V. Kudzma, A. Farid & J.L. Frank, *H. hortonii* (A.H. Sm. & Thiers) M. Kuo & B. Ortiz, *H. impositum* (Fr.) Šutara (typus), *H. indecorum* (Massee) G. Wu & Zhu L. Yang, *H. parvum* Mei Xiang Li, Zhu L. Yang & G. Wu, *H. rubropunctum* (Peck) Halling & B. Ortiz, *H. rugosum* G. Wu & Zhu L. Yang, *H. subglabripes* (Peck) Halling (Index Fungorum, accessed on 23 March 2023). *Hemileccinum* species share the following combination of characters: boletoid basidiomata, glabrous to subtomentose, smooth to rugose pileus surface, which turns violet with NH_3 vapours; tubes depressed around the stipe apex, pores at first light yellow to deep yellow becoming olive-yellow in age, concolorous with tubes, unchanging; olive spore deposit; central stipe, whose surface is always ornamented with scales concolorous with stipe, unchanging; pale yellow to light yellow context, unchanging; pileipellis a trichodermium with broad hyphae or an epithelium, sometime with filamentous terminal elements; pleurocystidia present, fusoid to lageniform; spores boletoid, subfusoid or ellipsoid in face view, smooth under light microscope, irregularly tiny warted and pinholed or rarely smooth under SEM; clamp connections absent (Šutara 2008; Halling et al. 2015; Wu et al. 2016; Index Fungorum 443:1, 2020; Kuo and Ortiz-Santana 2020; Farid et al. 2021; Li et al. 2021).

The first study of poroid mushrooms from Thailand was published in 1902, with descriptions of five new species, namely *Boletus lacunosus* Rostr. [current name: *Austroboletus rostrupii* (Syd. & P. Syd.) E. Horak], *Boletus costatus* Rostr., *Suillus changensis* Rostr. [current name: *Boletus changensis* (Rostr.) Sacc. & D. Sacc.], *Suillus hygrophanus* Rostr. [current name: *Boletus hygrophanus* (Rostr.) Sacc. & D. Sacc.], and *Suillus velatus* Rostr. [current name: *Veloporphyrillus velatus* (Rostr.) Y.C. Li & Zhu L. Yang] (Rostrup 1902; Saccardo and Saccardo 1905;

Horak 1980; Li et al. 2014). At that time, they were classified to belong to the Polyporaceae; however, later they were all moved to family Boletaceae. No new taxa in Boletaceae were described from Thailand during the following one hundred years. It is only in 2006 that again a new species, *Rhodactina incarnata* Zhu L. Yang, Trappe & Lumyong, was described from Chiang Mai Province, northern Thailand (Yang et al. 2006). In 2009, *Spongiforma thailandica* Desjardin, Manfr. Binder, Roekring & Flegel was described as a new genus and species from Nakorn Nayok Province, central Thailand (Desjardin et al. 2009). After that, molecular phylogenetic analyses have been widely used in Boletaceae taxonomy. Two more new Boletaceae genera including *Cacaoporus* Raspé & Vadthanarat and *Rubinosporus* Vadthanarat, Raspé & Lumyong were described from Chiang Mai Province, northern Thailand (Vadthanarat et al. 2019b, 2022). During that period, twenty-seven new species were also described from the country, among which nine belong in subfamily Xerocomoideae, namely *Heimioporus subcostatus* Vadthanarat, Raspé & Lumyong, *Phylloporus castanopsidis* M.A. Neves & Halling, *P. dimorphus* M.A. Neves & Halling, *P. infuscatus* M.A. Neves & Halling, *Phylloporus pusillus* Raspé, K.D. Hyde & Chuankid, *P. rubiginosus* M.A. Neves & Halling, *P. subrubeolus* Chuankid, K.D. Hyde & Raspé, *Rubinosporus auriporus* Vadthanarat, Raspé & Lumyong, *Xerocomus sisongkhramensis* Khamsuntorn, Pinruan & Luangsa-ard (Neves et al. 2012; Halling et al. 2014; Raspé et al. 2016; Vadthanarat et al. 2018; Chuankid et al. 2019; Vadthanarat et al. 2019a, 2019b, 2020; Chuankid et al. 2021; Raghoonundon et al. 2021; Vadthanarat et al. 2021; Tan et al. 2022; Vadthanarat et al. 2022).

In this study, several collections of boletes belonging to the subfamily Xerocomoideae were obtained from northern and northeastern Thailand. They were carefully studied based on morphology as well as family-wide and subfamily-wide phylogenetic analyses. Some of them were identified as a new *Hemileccinum* species. Some collections were identified as *X. sisongkhramensis* based on morphological characters and the megablast result of the ITS region. However, following multiple gene phylogenetic analyses based on four protein-coding gene (*atp6*, *cox3*, *tef1*, and *rpb2*), *X. sisongkhramensis* appeared phylogenetically distant from other *Xerocomus* species and distinct from existing genera in Boletaceae. Moreover, the detailed morphology did not fit any known Xerocomoideae genus. Therefore, *Rostrupomyces* is introduced to accommodate *X. sisongkhramensis*. Finally, a new *Hemileccinum* species is introduced with full descriptions and illustrations.

Materials and methods

Specimens collecting

Fresh basidiomata of boletes in subfamily Xerocomoideae were collected in Chiang Mai and Chiang Rai provinces in northern Thailand, and Ubon Ratchathani and Sisaket provinces in northeastern Thailand between 2015 and 2021. They were photographed in the field and then wrapped in aluminum foil for later description in the laboratory on the same day. The specimens were then dried in an electric drier at 45–50 °C. Examined specimens were deposited at MFU, BKF or CMUB herbaria.

Morphological study

Macroscopic descriptions were made based on the detailed field notes and photos of fresh basidiomata. Color codes were given based on Kornerup and Wanscher (1978). Macrochemical reactions (color reactions) were observed using aqueous solutions of 10% potassium hydroxide (KOH), and 28–30% NH_4OH . Microscopic structures were observed from dried specimens rehydrated in 5% KOH or 1% ammoniacal Congo red. For the measurements of microscopic features, a minimum of 50 basidiospores or 20 for other structures, were randomly chosen and measured under a Nikon Eclipse Ni compound microscope using NIS-Elements D version 5.10 software. The notation '[x/y/z]' represents the number of basidiospores 'x' measured from the number of basidiomata 'y' of the number of collections 'z'. The measurements of microscopic structures are presented in the following format (a–) b–c–d (–e), in which 'c' represents the average, 'b' is the 5th percentile, 'd' is the 95th percentile, and 'a' and 'e' the extreme values, shown in parentheses. *Q* is the length/width ratio. Sections of the pileipellis were cut radially, perpendicularly to the surface halfway between the centre and margin of pileus. Sections of stipeipellis were taken halfway along the stipe length (Li et al. 2011; Hosen et al. 2013; Li et al. 2014; Zhu et al. 2015). All line drawings of microscopic features were drawn by free hand using an Olympus compound microscope model CX41 with Olympus Camera Lucida model U-DA. For scanning electron microscopy, small fragments of dried hymenophore were mounted directly onto a SEM stub with double-sided carbon tape. The samples were coated with gold, examined and photographed using a TESCAN MIRA's 4th generation SEM.

DNA extraction, PCR amplification and DNA sequencing

Genomic DNA was extracted from tissue of dried specimen or fresh tissue preserved in CTAB, using a CTAB isolation procedure adapted from Doyle and Doyle (1990). Portions of the genes *atp6*, *cox3*, *rpb2*, and *tef1* were amplified by polymerase chain reaction (PCR). The primer pairs ATP6-1M40F/ATP6-2M (Raspé et al. 2016), COX3M1-F/ COX3M1-R (Vadthanarat et al. 2019b), bRPB2-6F/bRPB2-7.1R (Matheny 2005), and EF1-983F/EF1-2218R (Rehner and Buckley 2005) were used to amplify *atp6*, *cox3*, *rpb2*, and *tef1*, respectively. PCR products were purified by adding 1 U of exonuclease I and 0.5 U FastAP alkaline phosphatase (Thermo Scientific, St. Leon-Rot, Germany) and incubated at 37 °C for 1 h, followed by inactivation at 80 °C for 15 min. Standard Sanger sequencing was performed in both directions by Macrogen with PCR primers, except for *atp6*, for which universal primers M13F-pUC(-40) and M13F(-20) were used. For *tef1*, additional sequencing was performed with two internal primers, EF1-1577F and EF1-1567R (Rehner and Buckley 2005).

Alignment and phylogeny inference

The two reads of newly generated sequences were assembled in GENEIOUS Pro v. 6.0.6 (Biomatters) and blasted against GenBank database to check that they were not from unrelated contamination. For the Boletaceae-wide tree, the introns in *rpb2* and *tef1* were removed based on the amino acid sequence of

previously published sequences. The sequence datasets including the newly generated sequences and selected sequences representative of the whole family downloaded from GenBank, were separately aligned for each gene using MAFFT on the server accessed at <http://mafft.cbrc.jp/alignment/server/> (Kato and Standley 2013). Before combining the four gene partitions (*atp6*, *cox3*, *rpb2* exons + *tef1* exons), topological incongruence between the datasets was assessed using maximum likelihood (ML) on each of mitochondrial genes (*atp6* + *cox3*) dataset and nuclear genes (*rpb2* exons + *tef1* exons) dataset. Paired trees were examined for conflicts involving only nodes with ML bootstrap (BS) $\geq 70\%$. After that, the Maximum likelihood phylogenetic inference was performed using RAxML (Stamatakis 2006) on the CIPRES web portal (RAxML-HPC2 on XSEDE; Miller et al. 2009). The phylogenetic tree was inferred by a single partitioned analysis with four character sets (one for each gene), using the GTRCAT model with 25 categories. The outgroup consisted of two *Buchwaldoboletus* and seven *Chalciporus* species from subfamily Chalciporoideae, based on previously published phylogenies. Statistical support of clades was obtained with 1,000 rapid bootstrap replicates. For Bayesian Inference (BI), the best-fit model of substitution among those implementable in MrBayes was estimated separately for each region using jModel-test (Darriba et al. 2012) on the CIPRES portal, based on the Bayesian Information Criterion (BIC). The selected models were HKY+I+G for *atp6*, GTR+I+G for *cox3*, K80+I+G for *rpb2* exons, and SYM+I+G for *tef1* exons. Partitioned Bayesian analysis was performed on the CIPRES web portal (MrBayes on XSEDE; Ronquist et al. 2012). Two runs of five chains were run for 15,000,000 generations and sampled every 1,000 generations. At the end of the run, the average deviation of split frequencies was 0.008563. The PSRF values were equal or greater than 1, and ESS values were greater than 200 for all parameters. A total of 11,252 trees were used to construct a 50% majority rule consensus tree and calculate the Bayesian posterior probabilities (BPPs).

A second, Xerocomoideae-wide tree, was also inferred from sequences of selected taxa in Xerocomoideae. Sequences were also separately aligned for each of the genes using the MAFFT online software, with introns included. Then, the topological incongruence between the datasets was also assessed using ML on each gene of five character sets, *atp6*, *cox3*, *rpb2* exons, *tef1* exons, and the three introns of *tef1* + an intron of *rpb2*. Since there was no supported conflict, the ML phylogenetic tree was inferred by a single partitioned analysis with the five character sets (*atp6*, *cox3*, *rpb2* exons, *tef1* exons, and *rpb2* intron + *tef1* introns), using the same software and model that was used for family Boletaceae-wide phylogeny. Based on the latter, three *Hourangia*, three *Phylloporus*, and three *Xerocomus* species in the same subfamily Xerocomoideae were used as the outgroup. For BI, partitioned Bayesian analysis was performed with MrBayes 3.2.6 software for Windows. The selected models were GTR+I+G for *atp6* and *cox3*, K80+I+G for *rpb2* exons, and SYM+I+G for *tef1* exons, HKY+I+G intron of *rpb2* + introns of *tef1*. Two runs of five chains were sampled every 200 generations and stopped after 700,000 generations. At the end of the run, the average deviation of split frequencies was 0.007178. The PSRF values were equal or greater than 1, and ESS values were greater than 200 for all parameters. A total of 2,495 trees were used to construct a 50% majority rule consensus tree and calculate the BPPs.

Results

Phylogenetic analyses

A total of 39 sequences were newly generated in this study and deposited in GenBank. The ML phylograms from the mitochondrial and nuclear datasets were similar in topology without any supported conflict. The Boletaceae-wide, two-genome alignment contained 743 sequences comprising four genes (146 for *atp6*, 110 for *cox3*, 231 for *rpb2*, 256 for *tef1*) from 262 voucher specimens (Table 1) corresponding to 254 species, and was 2946 characters long (DOI: 10.6084/m9.figshare.23301077). ML and BI trees of the concatenated four-character set showed similar topologies without any supported conflicts (Bootstrap Support values, BS ≥ 70% and posterior probabilities, PP ≥ 0.90; Fig. 1). In the four-gene ML phylogram, the six subfamily clades were retrieved, namely the Austroboletoideae G. Wu & Zhu L. Yang, Boletoidae Singer, Chalciporoideae G. Wu & Zhu L. Yang, Leccinoideae G. Wu & Zhu L. Yang, Xerocomoideae, and Zangioideae G. Wu, Yan C. Li & Zhu L. Yang. The *Pulveroboletus* group introduced by Wu et al. (2014, 2016) was not monophyletic; however, the monophyly of each genus in this group was highly supported. All the *Xerocomus* (*Rostrupomyces*) *sisongkhramensis* collections included formed a highly supported (BS = 100%, PP = 1) monophyletic group, sister to *Rubinosporus* (BS = 99%, PP = 1) clustered in subfamily Xerocomoideae with high support (BS = 99%, PP = 1). The other selected *Xerocomus* species, including the type species *X. subtomentosus* (voucher VDKO 0987), formed another, distinct monophyletic group (BS = 89%, PP = 1), sister to *Phylloporus* (BS = 79%, PP = 1). The two genera also clustered in a supported clade together with *Hourangia* (BS = 100%, PP = 1). Regarding *Hemileccinum*, all selected species formed a highly supported clade (BS = 100%, PP = 1) consisting of fourteen species-level clades, including twelve known species, one new species from Thailand (this study), and one undescribed species from China. The new species *Hemileccinum inferius* clustered in a supported clade (BS = 76%, PP = 0.98) together with the American *H. hortonii*, the Chinese *H. rugosum*, and an undescribed *Hemileccinum* species from China (voucher HKAS53421).

Table 1. List of collections used for DNA analyses, with origin, GenBank accession numbers, and reference(s).

Species	Voucher	Origin	<i>atp6</i>	<i>cox3</i>	<i>rpb2</i>	<i>tef1</i>	Reference(s)
<i>Afroboletus</i> aff. <i>multijugus</i>	JD671	Burundi	MH614651	MH614794	MH614747	MH614700	Vadthanarat et al. (2019b)
<i>Afroboletus costatisporus</i>	ADK4644	Togo	KT823958	MH614795*	KT823991	KT824024	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Afroboletus luteolus</i>	ADK4844	Togo	MH614652	MH614796	MH614748	MH614701	Vadthanarat et al. (2019b)
<i>Amoenoboletus granulopunctatus</i>	HKAS 86007	China	–	–	MW560079	MZ741478	Wu et al. (2021)
<i>Amoenoboletus granulopunctatus</i>	HKAS 80250	China	–	–	MW560080	MW566746	Wu et al. (2021)
<i>Amylotrama banrockensis</i>	AD-C58672	Australia	–	–	–	MN413637	Lebel et al. (2022)
<i>Amylotrama clelandii</i>	MEL2432546	Australia	–	–	–	MN413630	Lebel et al. (2022)
<i>Anthracoporus cystidiatus</i>	HKAS55375	China	–	–	MT110410	KT990816*	Li and Yang (2021); Wu et al. (2016)*
<i>Anthracoporus holophaeus</i>	HKAS59407	China	–	–	KT990506	KT990888	Wu et al. (2016)
<i>Anthracoporus nigropurpureus</i>	HKAS52685	China	–	–	KT990459	KT990821	Wu et al. (2016)

Species	Voucher	Origin	<i>atp6</i>	<i>cox3</i>	<i>rpb2</i>	<i>tef1</i>	Reference(s)
<i>Aureoboletus auriflammeus</i>	CFMR:BOS-699	USA	–	–	MK766269	MK721060	Kuo and Ortiz-Santana (2020)
<i>Aureoboletus catenarius</i>	HKAS54467	China	–	–	KT990349	KT990711	Wu et al. (2016)
<i>Aureoboletus duplicatoporus</i>	HKAS50498	China	–	–	KF112754	KF112230	Wu et al. (2014)
<i>Aureoboletus formosus</i>	GDGM44441	China	–	–	KT291751	KT291744	Zhang et al. (2015)
<i>Aureoboletus gentilis</i>	ADK4865	Belgium	KT823961	MH614797*	KT823994	KT824027	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Aureoboletus glutinosus</i>	GDGM44477	China	–	–	MH700229	MH700205	Zhang et al. (2019)
<i>Aureoboletus innixus</i>	CFMR:BOS-544	USA	–	–	MK766270	MK721061	Kuo and Ortiz-Santana (2020)
<i>Aureoboletus moravicus</i>	VDK01120	Belgium	MG212528	MH614798*	MG212615	MG212573	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Aureoboletus nephrosporus</i>	HKAS74929	China	–	–	KT990358	KT990721	Wu et al. (2016)
<i>Aureoboletus pseudoauriporus</i>	JAB 80	USA	–	–	MW737471	MW737490	Farid et al. (2021)
<i>Aureoboletus raphanaceus</i>	GDGM 53127	China	–	–	MN549706	MN549676	Zhang et al. (2019)
<i>Aureoboletus singeri</i>	CFMR:BOS-468	Belize	–	–	MK766274	MK721065	Kuo and Ortiz-Santana (2020)
<i>Aureoboletus tenuis</i>	GDGM42601	China	–	–	KT291754	KT291745	Zhang et al. (2015)
<i>Aureoboletus thibetanus</i>	AFTOL-ID-450	China	DQ534600*	–	DQ366279	DQ029199	Binder and Hibbett (2006)*; Unpublished
<i>Aureoboletus tomentosus</i>	HKAS90216	China	–	–	KT990355	KT990717	Wu et al. (2016)
<i>Aureoboletus viscidipes</i>	HKAS77103	China	–	–	KT990360	KT990723	Wu et al. (2016)
<i>Aureoboletus viscosus</i>	OR0361	Thailand	MH614655	MH614801	MH614751	MH614704	Vadthanarat et al. (2019b)
<i>Australopilus palumanus</i>	REH-9433	Australia	–	–	MK766276	MK721067	Kuo and Ortiz-Santana (2020)
<i>Austroboletus</i> cf. <i>dictyotus</i>	OR0045	Thailand	KT823966	MH614802*	KT823999	KT824032	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Austroboletus</i> cf. <i>subvirens</i>	OR0573	Thailand	MH614656	MH614803	MH614752	MH614705	Vadthanarat et al. (2019b)
<i>Austroboletus olivaceoglutinosus</i>	HKAS57756	China	–	–	KF112764	KF112212	Wu et al. (2014)
<i>Baorangia major</i>	OR0209	Thailand	MG897421	MK372295*	MG897441	MG897431	Phookamsak et al. (2019); Vadthanarat et al. (2019b)*
<i>Baorangia pseudocalopus</i>	HKAS63607	China	–	–	KF112677	KF112167	Wu et al. (2014)
<i>Baorangia rufomaculata</i>	BOTH4144	USA	MG897415	MH614805*	MG897435	MG897425	Phookamsak et al. 2019; Vadthanarat et al. (2019b)*
<i>Binderoboletus segoi</i>	TWH8035	Guyana	OP358290	OP358307	–	–	This study
<i>Boletellus</i> aff. <i>ananas</i>	NY815459	Costa Rica	–	–	KF112760	KF112308	Wu et al. (2014)
<i>Boletellus</i> aff. <i>emodensis</i>	OR0061	Thailand	KT823970	MH614806*	KT824003	KT824036	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Boletellus ananas</i>	K(M)123769	Belize	MH614658	MH614807	MH614754	MH614707	Vadthanarat et al. (2019b)
<i>Boletellus areolatus</i>	TNS-F-61444 or BLT-7	Japan	–	AB989025	AB999754	–	Sata and Hattori (2015)
<i>Boletellus aurocontextus</i>	TNS-F-61501 or BLT-65	Japan	–	AB989037	AB999770	–	Sata and Hattori (2015)
<i>Boletellus emodensis</i>	TNS-F-61564 or BLT-128	Japan	–	AB989053	AB999782	–	Sata and Hattori (2015)
<i>Boletus aereus</i>	VDK01055	Belgium	MG212530	MH614809*	MG212617	MG212575	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Boletus albobrunnescens</i>	OR0131	Thailand	KT823973	MH614810*	KT824006	KT824039	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Boletus botryoides</i>	HKAS53403	China	–	–	KT990375	KT990738	Wu et al. (2016)
<i>Boletus edulis</i>	VDK00869	Belgium	MG212531	MH614811*	MG212618	MG212576	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Boletus rubriceps</i>	MICH:KUO-08150719	USA	–	–	MK766284	MK721076	Kuo and Ortiz-Santana (2020)
<i>Borofutus dhakanus</i>	OR0345	Thailand	MH614660	MH614814	MH614755	MH614709	Vadthanarat et al. (2019b)
<i>Buchwaldoboletus lignicola</i>	HKAS76674	China	–	–	KF112819	KF112277	Wu et al. (2014)
<i>Buchwaldoboletus lignicola</i>	VDK01140	Belgium	MH614661	MH614815	MH614756	MH614710	Vadthanarat et al. (2019b)

Species	Voucher	Origin	<i>atp6</i>	<i>cox3</i>	<i>rpb2</i>	<i>tef1</i>	Reference(s)
<i>Butyriboletus appendiculatus</i>	VDK00193b	Belgium	MG212537	MH614816*	MG212624	MG212582	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Butyriboletus</i> cf. <i>roseoflavus</i>	OR0230	China	KT823974	MH614819*	KT824007	KT824040	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Butyriboletus pseudoregius</i>	VDK00925	Belgium	MG212538	MH614817*	MG212625	MG212583	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Butyriboletus roseopurpureus</i>	BOTH4497	USA	MG897418	MH614818*	MG897438	MG897428	Phookamsak et al. (2019); Vadthanarat et al. (2019b)*
<i>Butyriboletus subsplendidus</i>	HKAS50444	China	–	–	KT990379	KT990742	Wu et al. (2016)
<i>Butyriboletus yicibus</i>	HKAS55413	China	–	–	KF112674	KF112157	Wu et al. (2014)
<i>Cacaoporus pallidicarneus</i>	SV0221	Thailand	MK372262	MK372299	MK372286	MK372273	Vadthanarat et al. (2019b)
<i>Cacaoporus tenebrosus</i>	SV0223	Thailand	MK372266	MK372303	MK372290	MK372277	Vadthanarat et al. (2019b)
<i>Caloboletus calopus</i>	ADK4087	Belgium	MG212539	MH614820	KP055030	KJ184566	Vadthanarat et al. (2018); Zhao et al. (2014a); Zhao et al. (2014b); Vadthanarat et al. (2019b)
<i>Caloboletus firmus</i>	BOS-372	Belize	–	–	MK766288	MK721080	Kuo and Ortiz-Santana (2020)
<i>Caloboletus inedulis</i>	BOTH3963	USA	MG897414	MH614821*	MG897434	MG897424	Phookamsak et al. (2019); Vadthanarat et al. (2019b)*
<i>Caloboletus radicans</i>	VDK01187	Belgium	MG212540	MH614822*	MG212626	MG212584	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Caloboletus yunnanensis</i>	HKAS69214	China	–	–	KT990396	KJ184568	Zhao et al. (2014a); Wu et al. (2016)
<i>Chalciporus</i> aff. <i>piperatus</i>	OR0586	Thailand	KT823976	MH614824*	KT824009	KT824042	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Chalciporus</i> aff. <i>rubinus</i>	OR0139	China	MH614663	–	MH614758	MH614712	Vadthanarat et al. (2019b)
<i>Chalciporus africanus</i>	JD517	Cameroon	KT823963	MH614825*	KT823996	KT824029	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Chalciporus piperatus</i>	VDK01063	Belgium	MH614664	MH614826	MH614759	MH614713	Vadthanarat et al. (2019b)
<i>Chalciporus rubinus</i>	AF2835	Belgium	KT823962	–	KT823995	KT824028	Raspé et al. (2016)
<i>Chalciporus</i> sp.	OR0363	Thailand	MH645586	MH645607	MH645602	MH645594	Vadthanarat et al. (2019b)
<i>Chalciporus</i> sp.	OR0373	Thailand	MH645587	MH645608	MH645603	MH645595	Vadthanarat et al. (2019b)
<i>Chamonixia brevicolumna</i>	DBG_F28707	USA	–	–	MK766291	MK721083	Kuo and Ortiz-Santana (2020)
<i>Chamonixia caespitosa</i>	OSC117571	USA	–	–	MK766293	MK721085	Kuo and Ortiz-Santana (2020)
<i>Chiuma virens</i>	OR0266	China	MG212541	MH614828*	MG212627	MG212585	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Chiuma viridula</i>	HKAS74928	China	–	–	KF112794	KF112273	Wu et al. (2014)
<i>Crocinoboletus</i> cf. <i>laetissimus</i>	OR0576	Thailand	KT823975	MH614833*	KT824008	KT824041	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Crocinoboletus rufoaureus</i>	HKAS53424	China	–	–	KF112710	KF112206	Wu et al. (2014)
<i>Cupreoboletus poikilochromus</i>	GS10070	Italy	–	–	KT157068	KT157072	Gelardi et al. (2015)
<i>Cyanoboletus brunneoruber</i>	OR0233	China	MG212542	MH614834*	MG212628	MG212586	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Cyanoboletus pulverulentus</i>	RW109	Belgium	KT823980	MH614835*	KT824013	KT824046	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Cyanoboletus sinopulverulentus</i>	HKAS59609	China	–	–	KF112700	KF112193	Wu et al. (2014)
<i>Erythrophylloporus aurantiacus</i>	REH7271	Costa Rica	MH614666	MH614829	MH614761	MH614715	Vadthanarat et al. (2019a)
<i>Erythrophylloporus fagicola</i>	Garay215	Mexico	MH614667	MH614830	MH614762	MH614716	Vadthanarat et al. (2019a)
<i>Erythrophylloporus paucicarpus</i>	OR1151	Thailand	MH614670	MH614831	MH614765	MH614719	Vadthanarat et al. (2019a)
<i>Erythrophylloporus suthepensis</i>	SV0236	Thailand	MH614672	MH614832	MH614767	MH614721	Vadthanarat et al. (2019a)
<i>Fistulinella prunicolor</i>	REH9880	Australia	MH614676	MH614840	MH614771	MH614725	Vadthanarat et al. (2019b)
<i>Harrya chromapes</i>	HKAS50527	China	–	–	KF112792	KF112270	Wu et al. (2014)
<i>Harrya moniliformis</i>	HKAS49627	China	–	–	KT990500	KT990881	Wu et al. (2016)
<i>Heimioporus conicus</i>	HKAS53451	China	–	–	KF112805	KF112226	Wu et al. (2016)

Species	Voucher	Origin	<i>atp6</i>	<i>cox3</i>	<i>rpb2</i>	<i>tef1</i>	Reference(s)
<i>Heimioporus australis</i>	REH9288	Australia	–	–	–	KP327703	Halling et al. (2015)
<i>Heimioporus cooloolae</i>	REH9817	Australia	–	–	–	KP327710	Halling et al. (2015)
<i>Heimioporus fruticola</i>	REH8962	Australia	–	–	–	KP327696	Halling et al. (2015)
<i>Heimioporus gaojiaocong</i>	HKAS80582	China	–	–	KT990409	KT990770	Wu et al. (2016)
<i>Heimioporus ivoryi</i>	REH8620	Costa Rica	–	–	–	KP327683	Halling et al. (2015)
<i>Heimioporus japonicus</i>	OR0114	Thailand	KT823971	–	KT824004	KT824037	Raspé et al. (2016)
<i>Heimioporus japonicus</i>	SV0016	Thailand	MT136776	–	MT136766	MT136771	Vadthanarat et al. (2020)
<i>Heimioporus mandarinus</i>	OR0218	Thailand	MG212546	–	MG212632	MG212590	Vadthanarat et al. (2018)
<i>Heimioporus subcostatus</i>	SV0235	Thailand	MT136780	–	MT136770	MT136775	Vadthanarat et al. (2020)
<i>Hemileccinum albidum</i>	KUN-HKAS81120	China	–	–	MZ936320	MZ936352	Li et. al. (2021)
<i>Hemileccinum inferius</i>	BR0260	Thailand	OP358291	–	OP358312	OP358319	This study
<i>Hemileccinum inferius</i>	SV0282	Thailand	OP358292	–	–	–	This study
<i>Hemileccinum brevisporum</i>	KUN-HKAS89150	China	–	–	MZ936328	MZ936362	Li et. al. (2021)
<i>Hemileccinum brevisporum</i>	HKAS59445	China	–	–	KT990414	KT990775	Wu et al. (2016)
<i>Hemileccinum depilatum</i>	AF2845	Belgium	MG212547	MH614843*	MG212633	MG212591	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Hemileccinum ferrugineipes</i>	KUN-HKAS115554	China	–	–	MZ936330	MZ973011	Li et. al. (2021)
<i>Hemileccinum floridanum</i>	AB16	USA	–	–	–	MW737481	Farid et al. (2021)
<i>Hemileccinum hortonii</i>	MICH:KUO-07050706	USA	–	–	MK766377	MK721175	Kuo and Ortiz-Santana (2020)
<i>Hemileccinum impolitum</i>	ADK4078	Belgium	MG212548	MH614844*	MG212634	MG212592	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Hemileccinum indecorum</i>	OR0863	Thailand	MH614677	MH614845	MH614772	MH614726	Vadthanarat et al. (2019b)
<i>Hemileccinum parvum</i>	KUN-HKAS115553	China	–	–	MZ936333	MZ973010	Li et. al. (2021)
<i>Hemileccinum rubropunctum</i>	REH-8501	USA	–	–	MK766327	MK721122	Kuo and Ortiz-Santana (2020)
<i>Hemileccinum rugosum</i>	HKAS84355	China	–	–	KT990413	KT990774	Wu et al. (2016)
<i>Hemileccinum</i> sp.	HKAS53421	China	–	–	KF112751	KF112235	Wu et al. (2014)
<i>Hemileccinum subglabripes</i>	MICH:KUO-07230802	USA	–	–	MK766300	MK721092	Kuo and Ortiz-Santana (2020)
<i>Hortiboletus amygdalinus</i>	HKAS54166	China	–	–	KT990416	KT990777	Wu et al. (2016)
<i>Hortiboletus campestris</i>	MICH:KUO-08240502	USA	–	–	MK766302	MK721094	Kuo and Ortiz-Santana (2020)
<i>Hortiboletus rubellus</i>	VDK00403	Belgium	MH614679	MH614847	MH614774	–	Vadthanarat et al. (2019b)
<i>Hortiboletus subpaludosus</i>	HKAS59608	China	–	–	KF112696	KF112185	Wu et al. (2014)
<i>Hourangia</i> cf. <i>pumila</i>	OR0762	Thailand	MH614680	MH614848	MH614775	MH614728	Vadthanarat et al. (2019b)
<i>Hourangia cheoi</i>	HKAS52269	China	–	–	KF112773	KF112286	Zhu et al. (2015)
<i>Hourangia microcarpa</i>	HKAS53378	China	–	–	KF112775	KF112300	Wu et al. (2014)
<i>Hourangia nigropunctata</i>	HKAS 57427	China	–	–	KP136978	KP136927	Zhu et al. (2015)
<i>Hymenoboletus luteopurpureus</i>	HKAS46334	China	–	–	KF112795	KF112271	Wu et al. (2014)
<i>Imleria badia</i>	VDK00709	Belgium	KT823983	MH614849*	KT824016	KT824049	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Imleria obscurebrunnea</i>	OR0263	China	MH614681	MH614850	MH614776	MH614729	Vadthanarat et al. (2019b)
<i>Imleria pallidus</i>	BOTH4356	USA	MH614659	MH614812	–	MH614708	Vadthanarat et al. (2019b)
<i>Indoporus squamulosus</i>	HKAS107153	China	–	–	MT110409	MT110335	Li and Yang (2021)
<i>Ionosporus longipes</i>	LEE1180	Malaysia	MT085461	–	MH712031*	MT085471	Chuankid et al. (2021); Khmelnitsky et al. (2019)
<i>Kaziboletus rufescens</i>	HKAS74706	Bangladesh	–	–	JQ928600	JQ928578	Hosen et al. (2021)

Species	Voucher	Origin	<i>atp6</i>	<i>cox3</i>	<i>rpb2</i>	<i>tef1</i>	Reference(s)
<i>Lanmaoa angustispora</i>	HKAS74752	China	–	–	KM605177	KM605154	Wu et al. (2015)
<i>Lanmaoa asiatica</i>	OR0228	China	MH614682	MH614851	MH614777	MH614730	Vadthanarat et al. (2019b)
<i>Lanmaoa carminipes</i>	BOTH4591	USA	MG897419	MH614852*	MG897439	MG897429	Phookamsak et al. (2019); Vadthanarat et al. (2019b)*
<i>Lanmaoa pallidrosea</i>	BOTH4432	USA	MG897417	MH614853*	MG897437	MG897427	Phookamsak et al. (2019); Vadthanarat et al. (2019b)*
<i>Lanmaoa sublurida</i>	Farid 1023	USA	–	–	MW737460	MW737485	Farid et al. (2021)
<i>Leccinellum</i> aff. <i>crocipodium</i>	HKAS76658	China	–	–	KF112728	KF112252	Wu et al. (2014)
<i>Leccinellum</i> aff. <i>griseum</i>	KPM-NC-0017832	Japan	KC552164	–	–	JN378450*	Unpublished; Orihara et al. (2012)*
<i>Leccinellum cremeum</i>	HKAS90639	China	–	–	KT990420	KT990781	Wu et al. (2016)
<i>Leccinum scabrum</i>	VDK00938	Belgium	MG212549	MH614858*	MG212635	MG212593	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Leccinum schistophilum</i>	VDK01128	Belgium	KT823989	MH614859*	KT824022	KT824055	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Leccinum variicolor</i>	VDK00844	Belgium	MG212550	MH614860*	MG212636	MG212594	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Leccinum versipelle</i>	KPM-NC-0017833	Scotland	KC552172	–	–	JN378454	Orihara et al. (2016); Orihara et al. (2012)
<i>Leccinum vulpinum</i>	KPM-NC-0017834	Scotland	KC552171	–	–	JN378456	Orihara et al. (2016); Orihara et al. (2012)
<i>Mucilopilus castaneiceps</i>	HKAS75045	China	–	–	KF112735	KF112211	Wu et al. (2014)
<i>Mucilopilus paracastaneiceps</i>	HKAS50338	China	–	–	KT990391	KT990755	Wu et al. (2016)
<i>Mucilopilus ruber</i>	HKAS84555	China	–	–	MT110436	MT110364	Li and Yang (2021)
<i>Mycoamaranthus cambodgensis</i>	SV0197	Thailand	MZ355900	MZ355909	–	–	Vadthanarat et al. (2022)
<i>Neoboletus brunneissimus</i>	OR0249	China	MG212551	MH614861*	MG212637	MG212595	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Neoboletus ferrugineus</i>	HKAS77718	China	–	–	KT990431	KT990789	Wu et al. (2016)
<i>Neoboletus flavidus</i>	HKAS59443	China	–	–	KU974144	KU974136	Wu et al. (2016)
<i>Neoboletus hainanensis</i>	HKAS59469	China	–	–	KF112669	KF112175	Wu et al. (2014)
<i>Neoboletus junquilleus</i>	AF2922	France	MG212552	MH614862*	MG212638	MG212596	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Neoboletus magnificus</i>	HKAS74939	China	–	–	KF112653	KF112148	Wu et al. (2014)
<i>Neoboletus obscureumbrinus</i>	OR0553	Thailand	MK372271	–	MK372294	MK372282	Vadthanarat et al. (2019b)
<i>Neoboletus tomentulosus</i>	HKAS53369	China	–	–	KF112659	KF112154	Wu et al. (2014)
<i>Neoboletus erythropus</i>	VDK00690	Belgium	KT823982	MH614864*	KT824015	KT824048	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Octaviania asterosperma</i>	AQUI3899	Italy	KC552159	–	–	KC552093	Orihara et al. (2016)
<i>Octaviania cyanescens</i>	PNW-FUNGI-5603	USA	KC552160	–	–	JN378438	Orihara et al. (2016); Orihara et al. (2012)
<i>Octaviania tasmanica</i>	MEL2128484	Australia	KC552157	–	–	JN378437	Orihara et al. (2016); Orihara et al. (2012)
<i>Octaviania zelleri</i>	MES270	USA	KC552161	–	–	JN378440	Orihara et al. (2016); Orihara et al. (2012)
<i>Parvixerocomus pseudoaokii</i>	OR0155	China	MG212553	MH614865	MG212597	MG212597	Vadthanarat et al. (2019b)
<i>Paxilloboletus latisporus</i>	ADK5072	Congo	–	–	MZ707870	MZ707866	Badou et al. (2022)
<i>Paxilloboletus africanus</i>	SAB0716	Guinea	–	–	MZ707869	MZ707865	Badou et al. (2022)
<i>Phylloporus bellus</i>	OR0473	China	MH580778	MH614866*	MH580818	MH580798	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus brunneiceps</i>	OR0050	Thailand	KT823968	MH614867*	KT824001	KT824034	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Phylloporus castanopsidis</i>	OR0052	Thailand	KT823969	MH614868*	KT824002	KT824035	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Phylloporus maculatus</i>	OR0285	China	MH580780	–	MH580820	MH580800	Chuankid et al. (2019)

Species	Voucher	Origin	atp6	cox3	rpb2	tef1	Reference(s)
<i>Phylloporus pachycystidiatus</i>	HKAS53422	China	–	–	KF112777	KF112288	Wu et al. (2014)
<i>Phylloporus pelletieri</i>	WU18746	Austria	MH580781	MH614869*	MH580821	MH580801	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus pusillus</i>	OR1158	Thailand	MH580783	MH614870*	MH580823	MH580803	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus rhodoxanthus</i>	WU17978	Austria	MH580785	MH614871*	MH580824	MH580805	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus rubeolus</i>	OR0251	China	MH580786	MH614872*	MH580825	MH580806	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus rubiginosus</i>	OR0169	China	MH580788	MH614873*	MH580827	MH580808	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus rubrosquamosus</i>	HKAS52552	China	–	–	KF112780	KF112289	Wu et al. (2014)
<i>Phylloporus scabripes</i>	CFMR:BOS-621	Belize	–	–	MK766359	MK721156	Kuo and Ortiz-Santana (2020)
<i>Phylloporus subbacillisporus</i>	OR0436	China	MH580792	MH614875*	MH580831	MH580812	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus subrubeolus</i>	BC022	Thailand	MH580793	MH614876*	MH580832	MH580813	Chuankid et al. (2019); Vadthanarat et al. (2019b)*
<i>Phylloporus yunnanensis</i>	OR0448	China	MG212554	MH614877*	MG212640	MG212598	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Porphyrellus castaneus</i>	OR0241	China	MG212555	MH614878*	MG212641	MG212599	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Porphyrellus porphyrosporus</i>	MB97 023	Germany	DQ534609	–	GU187800	GU187734	Binder and Hibbett (2006); Binder et al. (2010)
<i>Pseudoaustroboletus valens</i>	HKAS82644	China	–	–	MT110431	MT110359	Li and Yang (2021)
<i>Pulchroboletus sclerotiorum</i>	FLAS F 60333	USA	–	–	MF614169	MF614167	Crous et al. (2019)
<i>Pulchroboletus sclerotiorum</i>	FLAS F 60334	USA	–	–	MF614164	MF614165	Crous et al. (2019)
<i>Pulveroboletus</i> aff. <i>ravenelii</i>	ADK4360	Togo	KT823957	MH614882*	KT823990	KT824023	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Pulveroboletus</i> aff. <i>ravenelii</i>	ADK4650	Togo	KT823959	MH614883*	KT823992	KT824025	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Pulveroboletus brunneopunctatus</i>	HKAS55369	China	–	–	KT990455	KT990814	Wu et al. (2016)
<i>Pulveroboletus fragrans</i>	OR0673	Thailand	KT823977	MH614884*	KT824010	KT824043	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Pulveroboletus ravenelii</i>	REH2565	USA	KU665635	MH614885*	KU665637	KU665636	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Retiboletus</i> aff. <i>nigerrimus</i>	OR0049	Thailand	KT823967	MH614886*	KT824000	KT824033	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Retiboletus brevibasidiatus</i>	OR0570	Thailand	MT085469	–	MT085479	MT085476	Chuankid et al. (2021)
<i>Retiboletus brunneolus</i>	HKAS52680	China	–	–	KF112690	KF112179	Wu et al. (2014)
<i>Retiboletus fuscus</i>	OR0231	China	MG212556	MH614887*	MG212642	MG212600	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Retiboletus griseus</i>	MB03 079	USA	KT823964	MH614888*	KT823997	KT824030	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Retiboletus kauffmanii</i>	OR0278	China	MG212557	MH614889*	MG212643	MG212601	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Retiboletus nigerrimus</i>	HKAS53418	China	–	–	KT990462	KT990824	Wu et al. (2016)
<i>Rhodactina himalayensis</i>	CMU25117	Thailand	MG212558	–	–	MG212602, MG212603	Vadthanarat et al. (2018)
<i>Rhodactina rostratispora</i>	SV0170	Thailand	MG212560	–	MG212645	MG212605	Vadthanarat et al. (2018)
<i>Rossbeevera cryptocyanea</i>	KPM-NC17843	Japan	KT581441	–	–	KC552072	Orihara et al. (2016)
<i>Rossbeevera eucyanea</i>	TNS-F-36986	Japan	KC552115	–	–	KC552068	Orihara et al. (2016)
<i>Rossbeevera griseovelutina</i>	TNS-F-36989	Japan	KC552124	–	–	KC552076	Orihara et al. (2016)
<i>Rossbeevera pachydermis</i>	KPM-NC23336	New Zealand	KJ001064	–	–	KP222912	Orihara et al. (2016)
<i>Rossbeevera vittatispora</i>	TO-AUS-72	Australia	KC552108	–	–	KC552065	Orihara et al. (2016)

Species	Voucher	Origin	atp6	cox3	rpb2	tef1	Reference(s)
<i>Rostrupomyces sisongkhramensis</i>	BR0311	Thailand	OP358293	–	OP358313	OP358320	This study
<i>Rostrupomyces sisongkhramensis</i>	BR0313	Thailand	OP358294	–	OP358314	OP358321	This study
<i>Rostrupomyces sisongkhramensis</i>	BR0368	Thailand	OP358295	–	–	–	This study
<i>Rostrupomyces sisongkhramensis</i>	BR0371	Thailand	OP358296	–	–	OP358322	This study
<i>Rostrupomyces sisongkhramensis</i>	OR0915	Thailand	OP358297	–	–	–	This study
<i>Rostrupomyces sisongkhramensis</i>	OR0918	Thailand	OP358298	–	–	–	This study
<i>Rostrupomyces sisongkhramensis</i>	OR0919	Thailand	OP358299	OP358308	OP358315	OP358323	This study
<i>Rostrupomyces sisongkhramensis</i>	OR1004	Thailand	OP358300	–	–	–	This study
<i>Rostrupomyces sisongkhramensis</i>	OR1059	Thailand	OP358301	–	–	–	This study
<i>Rostrupomyces sisongkhramensis</i>	OR1392	Thailand	OP358302	–	–	–	This study
<i>Rostrupomyces sisongkhramensis</i>	OR1399	Thailand	OP358303	–	–	–	This study
<i>Rostrupomyces sisongkhramensis</i>	SV0155	Thailand	OP358304	OP358309	OP358316	OP358324	This study
<i>Rostrupomyces sisongkhramensis</i>	SV0219	Thailand	OP358305	OP358310	OP358317	OP358325	This study
<i>Rostrupomyces sisongkhramensis</i>	SV0225	Thailand	OP358306	OP358311	OP358318	OP358326	This study
<i>Royoungia rubina</i>	HKAS53379	China	–	–	KF112796	KF112274	Wu et al. (2014)
<i>Rubinosporus auriporus</i>	SV0101	Thailand	MZ355897	MZ355906	MZ355904	MZ355902	Vadthanarat et al. (2022)
<i>Rubinosporus auriporus</i>	SV0090	Thailand	MZ355896	MZ355905	MZ355903	MZ355901	Vadthanarat et al. (2022)
<i>Rubroboletus legaliae</i>	VDKO0936	Belgium	KT823985	MH614890*	KT824018	KT824051	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Rubroboletus rhodosanguineus</i>	BOTH4263	USA	MG897416	MH614891*	MG897436	MG897426	Phookamsak et al. (2019); Vadthanarat et al. (2019b)*
<i>Rubroboletus rhodoxanthus</i>	HKAS84879	China	–	–	KT990468	KT990831	Wu et al. (2016)
<i>Rubroboletus satanas</i>	VDKO0968	Belgium	KT823986	MH614892*	KT824019	KT824052	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Rugiboletus andinus</i>	REH-7705	Costa Rica	–	–	MK766316	MK721111	Kuo and Ortiz-Santana (2020)
<i>Rugiboletus brunneiporus</i>	HKAS83209	China	–	–	KM605168	KM605144	Wu et al. (2015)
<i>Rugiboletus extremiorientalis</i>	OR0406	Thailand	MG212562	MH614893*	MG212647	MG212607	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Singerocomus inundabilis</i>	TWH9199	Guyana	MH645588	MH645609	LC043089*	MH645596	Henkel et al. (2016)*; Vadthanarat et al. (2019b)
<i>Singerocomus rubriflavus</i>	TWH9585	Guyana	MH645589	MH645610	–	MH645597	Vadthanarat et al. (2019b)
<i>Spongiforma thailandica</i>	DED7873	Thailand	MG212563	MH614894**	MG212648	KF030436*	Nuhn et al. (2013)*; Vadthanarat et al. (2018); Vadthanarat et al. (2019b)**
<i>Spongispora temasekensis</i>	SING 0206334	Singapore	–	–	MG674378	MG674377	Wu et al. (2018)
<i>Spongispora temasekensis</i>	ACMF5	Singapore	MZ803018	–	MZ824748	MZ803023	Raghoonundon et al. (2021)
<i>Strobilomyces atosquamosus</i>	HKAS55368	China	–	–	KT990476	KT990839	Wu et al. (2016)
<i>Strobilomyces echinocephalus</i>	OR0243	China	MG212564	–	MG212649	MG212608	Vadthanarat et al. (2018)
<i>Strobilomyces floccopus</i>	RW103	Belgium	KT823978	MH614895*	KT824011	KT824044	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Strobilomyces mirandus</i>	OR0115	Thailand	KT823972	MH614896*	KT824005	KT824038	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Strobilomyces verruculosus</i>	HKAS55389	China	–	–	KF112813	KF112259	Wu et al. (2014)
<i>Suillellus luridus</i>	VDKO0241b	Belgium	KT823981	MH614901*	KT824014	KT824047	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Suillellus queletii</i>	VDKO1185	Belgium	MH645590	MH645611	MH645604	MH645598	Vadthanarat et al. (2019b)

Species	Voucher	Origin	atp6	cox3	rpb2	tef1	Reference(s)
<i>Suillellus subamygdalinus</i>	HKAS57262	China	–	–	KF112660	KF112174	Wu et al. (2014)
<i>Sutorius australiensis</i>	REH9441	Australia	MG212567	MK386576**	MG212652	JQ327032*	Halling et al. (2012)*; Vadthanarat et al. (2018); Vadthanarat et al. (2019b)**
<i>Sutorius eximius</i>	REH9400	USA	MG212568	MH614902**	MG212653	JQ327029*	Halling et al. (2012)*; Vadthanarat et al. (2018); Vadthanarat et al. (2019b)**
<i>Sutorius pachypus</i>	OR0411	Thailand	MN067465	–	MN067500	MN067484	Vadthanarat et al. (2021)
<i>Sutorius pseudotylopilus</i>	OR0378B	Thailand	MH614692	MH614903	MH614787	MH614740	Vadthanarat et al. (2019b)
<i>Sutorius rubinus</i>	OR0379	Thailand	MH614693	MH614904	MH614788	MH614741	Vadthanarat et al. (2019b)
<i>Sutorius ubonensis</i>	SV0032	Thailand	MN067472	–	MN067507	MN067491	Vadthanarat et al. (2021)
<i>Tengioboletus glutinosus</i>	HKAS53425	China	–	–	KF112800	KF112204	Wu et al. (2014)
<i>Tengioboletus reticulatus</i>	HKAS53426	China	–	–	KF112828	KF112313	Wu et al. (2014)
<i>Turmalinea persicina</i>	KPM-NC18001	Japan	KC552130	–	–	KC552082	Orihara et al. (2016)
<i>Turmalinea yuwanensis</i>	KPM-NC18011	Japan	KC552138	–	–	KC552089	Orihara et al. (2016)
<i>Tylocinum griseolum</i>	HKAS50281	China	–	–	KF112730	KF112284	Wu et al. (2014)
<i>Tylopilus atripurpureus</i>	HKAS50208	China	–	–	KF112799	KF112283	Wu et al. (2014)
<i>Tylopilus felleus</i>	VDK00992	Belgium	KT823987	MH614906*	KT824020	KT824053	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Tylopilus ferrugineus</i>	BOTH3639	USA	MH614694	MH614907	MH614789	MH614742	Vadthanarat et al. (2019b)
<i>Tylopilus otsuensis</i>	HKAS53401	China	–	–	KF112797	KF112224	Wu et al. (2014)
<i>Tylopilus vinaceipallidus</i>	OR0137	China	MG212571	MH614912*	MG212656	MG212613	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Tylopilus violaceobrunneus</i>	HKAS89443	China	–	–	KT990504	KT990886	Wu et al. (2016)
<i>Veloboletus limbatus</i>	REH9228	Australia	MT747398	–	MT747397	MN413636	Crous et al. (2019)
<i>Veloporphyrellus conicus</i>	REH8510	Belize	MH614698	MH614913	MH614792	MH614745	Vadthanarat et al. (2019b)
<i>Veloporphyrellus gracilioides</i>	HKAS53590	China	–	–	KF112734	KF112210	Wu et al. (2014)
<i>Veloporphyrellus pseudovelatus</i>	HKAS59444	China	JX984519	–	–	JX984553	Li et al. (2014)
<i>Veloporphyrellus velatus</i>	HKAS63668	China	JX984523	–	–	JX984554	Li et al. (2014)
<i>Xanthoconium affine</i>	NY00815399	USA	–	–	KT990486	KT990850	Wu et al. (2016)
<i>Xanthoconium purpureum</i>	MICH:KUO-07061405	USA	–	–	MK766372	MK721170	Kuo and Ortiz-Santana (2020)
<i>Xanthoconium sinense</i>	HKAS77651	China	–	–	KT990488	KT990853	Wu et al. (2016)
<i>Xerocomellus chrysenteron</i>	VDK00821	Belgium	KT823984	MH614914*	KT824017	KT824050	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Xerocomellus cisalpinus</i>	ADK4864	Belgium	KT823960	MH614915*	KT823993	KT824026	Raspé et al. (2016); Vadthanarat et al. (2019b)*
<i>Xerocomellus communis</i>	HKAS50467	China	–	–	KT990494	KT990858	Wu et al. (2016)
<i>Xerocomellus ripariellus</i>	VDK00404	Belgium	MH614699	MH614916	MH614793	MH614746	Vadthanarat et al. (2019b)
<i>Xerocomus ferrugineus</i>	CFMR:BOS-545	USA	–	–	MK766375	MK721173	Kuo and Ortiz-Santana (2020)
<i>Xerocomus fulvipes</i>	HKAS76666	China	–	–	KF112789	KF112292	Wu et al. (2014)
<i>Xerocomus magniporus</i>	HKAS58000	China	–	–	KF112781	KF112293	Wu et al. (2014)
<i>Xerocomus rugosellus</i>	HKAS58865	China	–	–	KF112784	KF112294	Wu et al. (2014)
<i>Xerocomus spadiceus</i> var. <i>gracilis</i>	MICH:KUO-07080702	USA	–	–	MK766378	MK721176	Kuo and Ortiz-Santana (2020)
<i>Xerocomus subtomentosus</i>	VDK00987	Belgium	MG212572	MH614919*	MG212657	MG212614	Vadthanarat et al. (2018); Vadthanarat et al. (2019b)*
<i>Xerocomus tenax</i>	MICH:KUO-08241404	USA	–	–	MK766379	MK721177	Kuo and Ortiz-Santana (2020)
<i>Zangia citrina</i>	HKAS52684	China	HQ326850	–	–	HQ326872	Li et al. (2011)
<i>Zangia olivaceobrunnea</i>	HKAS52272	China	HQ326857	–	–	HQ326876	Li et al. (2011)
<i>Zangia roseola</i>	HKAS51137	China	HQ326858	–	–	HQ326877	Li et al. (2011)

For the subfamily Xerocomoideae-wide phylogeny, no supported topological incongruence between the character sets was detected. Then, the Xerocomoideae-wide phylogeny was inferred based on the alignment containing 155 sequences of four genes (22 for *atp6*, 20 for *cox3*, 53 for *rpb2*, 60 for *tef1*) from 60 voucher specimens corresponding to 55 taxa, and was 3,161 characters long (DOI: 10.6084/m9.figshare.23301077). The ML and BI tree topologies

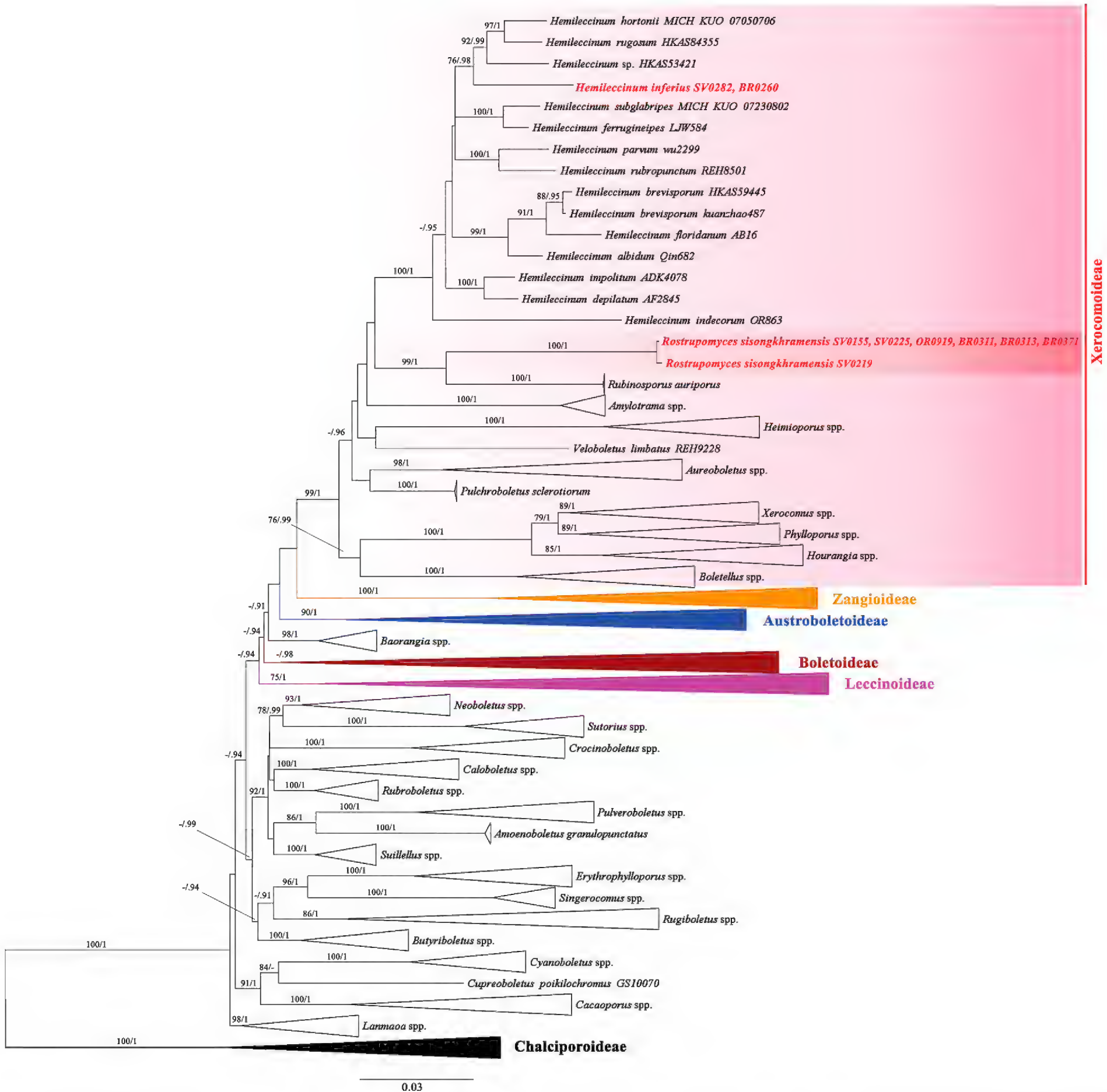


Figure 1. Boletaceae-wide Maximum Likelihood phylogenetic tree inferred from the four-gene dataset (*atp6*, *cox3*, *rpb2*, and *tef1*) (introns excluded), showing the position of the new genus *Rostrupomyces* in Xerocomoideae. Bootstrap support values (BS $\geq 70\%$) and the corresponding Bayesian posterior probabilities (PP ≥ 0.90) are shown above the supported branches. The two *Buchwaldoboletus* and seven *Chalciporus* species (subfamily Chalciporoideae) were used as outgroup. All taxa belonging to subfamilies Austroboletioideae, Boletioideae, Chalciporoideae, Leccinoideae, and Zangioideae were collapsed into subfamily clades. All generic clades in subfamily Xerocomoideae (excluding *Hemileccinum* and *Rostrupomyces*) and *Pulveroboletus* group with high supports, were also collapsed.

of the concatenated five-character-set alignment were similar without any supported conflict (Fig. 2). The Xerocomoideae-wide ML tree also showed a similar topology to the Boletaceae-wide tree. However, in this subfamily Xerocomoideae-wide tree, the support of the clade consisting of the new species *Hemileccinum inferius*, *H. hortonii*, *H. rugosum*, and an undescribed *Hemileccinum* species, was lower (BS = 53%, PP = 0.71) than in the Boletaceae-wide ML tree.

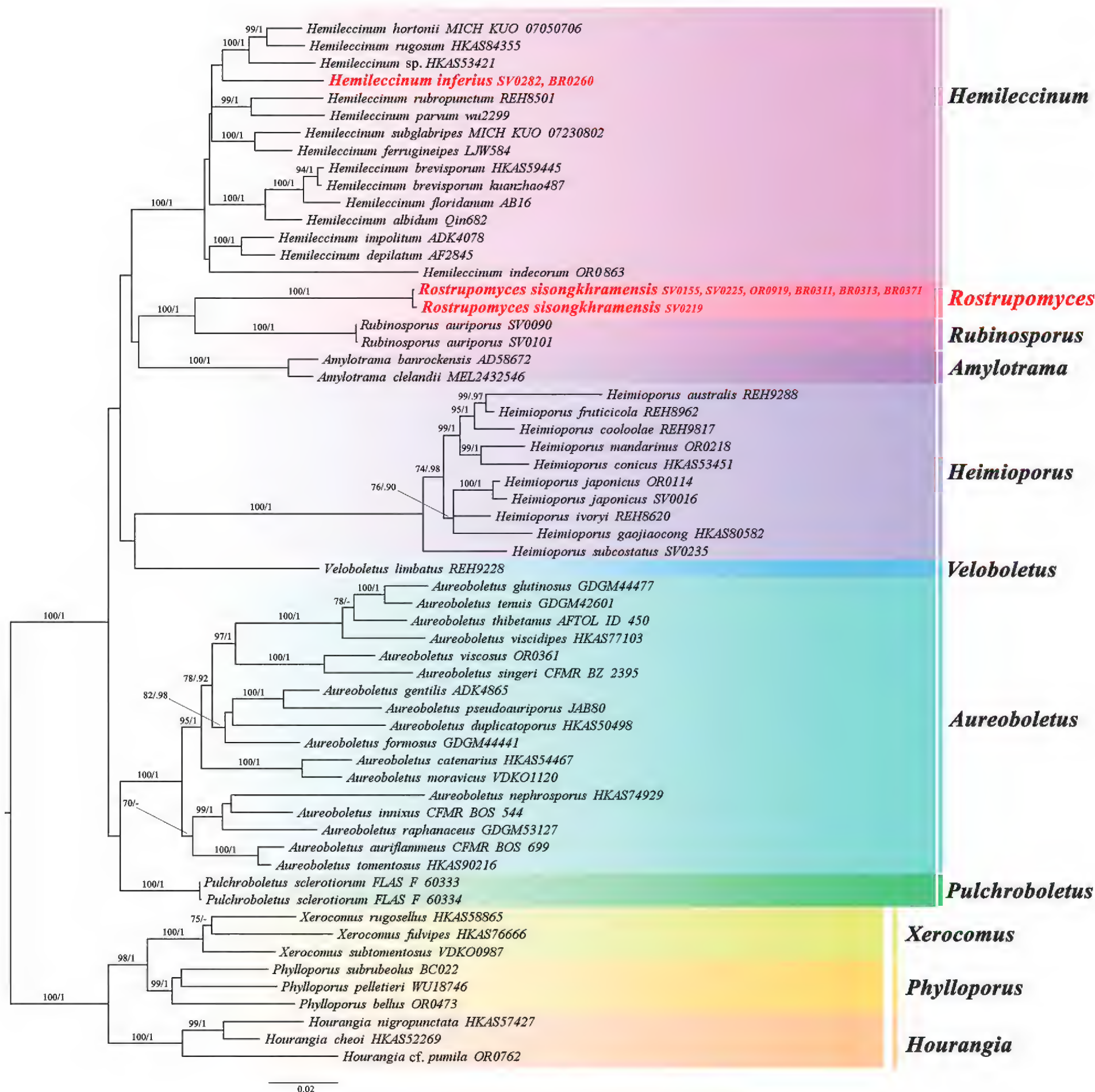


Figure 2. Xerocomoideae-wide phylogenetic tree inferred from the four-gene dataset (*atp6*, *cox3*, *rpb2*, and *tef1*) (introns included), including new genus *Rostrupomyces* and selected Xerocomoideae using Maximum Likelihood and Bayesian Inference methods (ML tree is presented). The three *Hourangia*, three *Phylloporus*, and three *Xerocomus* species in Xerocomoideae were used as outgroup. Bootstrap support values (BS \geq 70%) and posterior probabilities (PP \geq 0.90) are shown above the supported branches.

Taxonomy

Rostrupomyces Vadthanarat & Raspé, gen. nov.

MycoBank No: 849050

Etymology. Named in honor of Frederik Georg Emil Rostrup (1831–1907), Danish botanist, mycologist, and plant pathologist, celebrating the 120 years of his describing the first new species of Boletaceae from Thailand in 1902.

Diagnosis. Differs from other genera in Boletaceae by the following combination of characters: rugulose to subrugulose pileus surface, white pore when young becoming grayish yellow in age, subscabrous stipe surface with scattered granulate squamules, white basal mycelium, unchanging color in any parts, yellowish brown spore print, and broadly ellipsoid to ellipsoid, smooth basidiospores.

Description. *Basidiomata* stipitate-pileate. *Pileus* convex then plano-convex to plane; *surface* at first rugulose then subrugulose in age, finely tomentose to tomentose, dark brown to reddish brown, becoming light brown to brown to grayish orange, unchanging when bruised; *context* off-white then yellowish to dull pale orange in age, unchanging when cut. *Stipe* central, terete, cylindrical; *surface* subscabrous, yellowish white to pale yellow to orange white, with scattered brown to dark brown to reddish brown granulate squamules, unchanging when bruised; *basal mycelium* white; *context* solid, white becoming off-white to yellowish white in age, unchanging when cut. *Hymenophore* tubulate, slightly depressed to depressed around the stipe. *Tubes* pale yellow then grayish yellow, separable from the pileus context, unchanging when cut. *Pores* roundish then subangular to angular with age; when young white then yellowish white becoming grayish yellow, unchanging when touched. *Spore print* yellowish brown. *Basidiospores* ellipsoid to broadly ellipsoid, thin-walled, smooth under light microscope and SEM. *Basidia* 4-spored, clavate without basal clamp connection. *Cheilo- and pleurocystidia* narrowly fusiform to fusiform or narrowly utriform, thin-walled. *Pileipellis* an intricate trichoderm, made of moderately interwoven to loosely interwoven, thin-walled hyphae. *Stipitipellis* arranged parallel to the surface of the stipe, composed of moderately interwoven, thin-walled hyphae, with scattered groups of rising cells to clusters of narrowly clavate to clavate cells. *Clamp connections* not seen in any tissue.

Typus generis. *Rostrupomyces sisongkhramensis* (Khamsuntorn, Pinruan & Luangsa-ard) Vadthanarat, Raghoonundon & Raspé.

Distribution. Currently known only from northern and northeastern Thailand.

Notes. *Rostrupomyces* can be morphologically separated from *Xerocomus* by the different shape and surface of basidiospores, which are ellipsoid to broadly ellipsoid with smooth under light microscope and SEM in the new genus, whereas *Xerocomus* produce more or less oblong to fusiform basidiospores, usually with bacillate surface under SEM (Wu et. al. 2016). *Rostrupomyces* also produces yellowish brown spore print, whereas *Xerocomus* produces olive-brown spore print. Moreover, color change upon bruising does not occur in any part of *Rostrupomyces* basidiomes, whereas context and hymenophore of *Xerocomus* always turn more or less bluish to blue when bruised or cut (Wu et. al. 2016). The most resembling genus, *Hemileccinum*, shares some similar characters including rugulose to subrugulose pileus surface, yellow hymeno-

phore which is depressed around the stipe apex, subscabrous stipe surface (less so in *Hemileccinum*), white basal mycelium, mostly unchanging color in any parts. However, *Rostrupomyces* can be morphologically distinguished from *Hemileccinum* by the differences in spore print color, and in the shape and surface of basidiospores. *Rostrupomyces* produces yellowish brown spore print, broadly ellipsoid to ellipsoid basidiospores with smooth surface under light microscope and SEM. *Hemileccinum* produces olive-brown spore prints, boletoid basidiospores that are smooth under light microscope, but ornamented with irregular warts and pinholes under SEM. Also, the pore surface of *Rostrupomyces* is white in young basidiomata and becomes pale yellow when mature whereas in *Hemileccinum*, the pore surface is yellow in all stages (Šutara 2008; Wu et al. 2016; Farid et al. 2021; Li et al. 2021).

***Rostrupomyces sisongkhramensis* (Khamsuntorn, Pinruan & Luangsa-ard)
Vadthanarat, Raghoonundon & Raspé, comb. nov.**

Figs 3, 4, 5A–B

MycoBank No: 851393

Xerocomus sisongkhramensis Khamsuntorn, Pinruan & Luangsa-ard. Basionym.

Diagnosis. *Rostrupomyces sisongkhramensis* is characterised by having dark to reddish brown, becoming brown to grayish orange pileus, with rugulose to subrugulose, finely tomentose to tomentose surface; yellowish to orange white, subscabrous, longitudinally fissurate stipe surface, with moderately scattered brown to dark brown to reddish brown granulate squamules; yellow hymenophore; unchanging color in any parts; yellowish brown spore print; and broadly ellipsoid to ellipsoid smooth basidiospores.

Description. *Basidiomata* medium-sized. **Pileus** 37–94(118) mm in diameter, convex at first then plano-convex to plane, sometimes with sub-depressed at the centre; **margin** inflexed at first then deflexed in age, exact or slightly exceeding (up to 1 mm); **surface** at first rugulose especially near the margin then subrugulose in age, dull, dry to moist, finely tomentose to tomentose covered with greenish yellow (3A3–4, 3B4) matted hyphae at places (especially when young), at first dark brown to reddish brown (6–8F4–8), becoming light brown to brown to grayish orange (6D/E5–6, 5B4–5) on light yellow to brownish orange (4A3–5, 5C4) background in age, gradually paler to the margin, unchanging when bruised; **context** (3)5–10(14) mm thick half-way to the margin, at first firm then soft in age, color distribution even, at first off-white, slightly brownish (7D/E4–5) near the pileipellis, then yellowish to orange white (4–5A2) or occasionally yellowish (3A3–4) above the hymenium especially in age, unchanging when cut. **Stipe** (33)41–97(108) × 6(7)–19(20) mm, central, terete, usually cylindrical for the most part but often with wider base, rarely club-shaped; **surface** subscabrous longitudinally fissurate, slightly shiny, yellowish white to pale yellow to orange white (3A3 to 4A2 to 5A2), occasionally pale yellow (3A3–4) near the cap, with moderately scattered brown to dark brown to reddish brown (7D/E/F4–7) granulate squamules, unchanging when bruised; **basal mycelium** little developed, white (1A1); **context** solid, firm, at first white (1A1) becoming off-white to yellowish white (4A2) occasionally pale yellow (3A3–4) especially

in the above part near the stipe surface in age, yellowish to orange gray (4–5B2–3) virgate at places, unchanging when cut. **Hymenophore** tubulate, slightly depressed to depressed around the stipe, with slightly decurrent tooth, sometimes almost free, mostly segmentiform to subventricose. **Tubes** (3)4–13 mm long half-way to the margin, at first pale yellow (4A3) then grayish yellow (4B3)



Figure 3. Fresh basidiomata of *Rostrupomyces sisongkhamensis* **A** OR0915 **B** OR0919 **C** OR1004 **D** SV0155, white pores surface in young basidioma (white arrow) **E** SV0219 **F** SV0225. Scale bars: 1 cm (**A–F**).

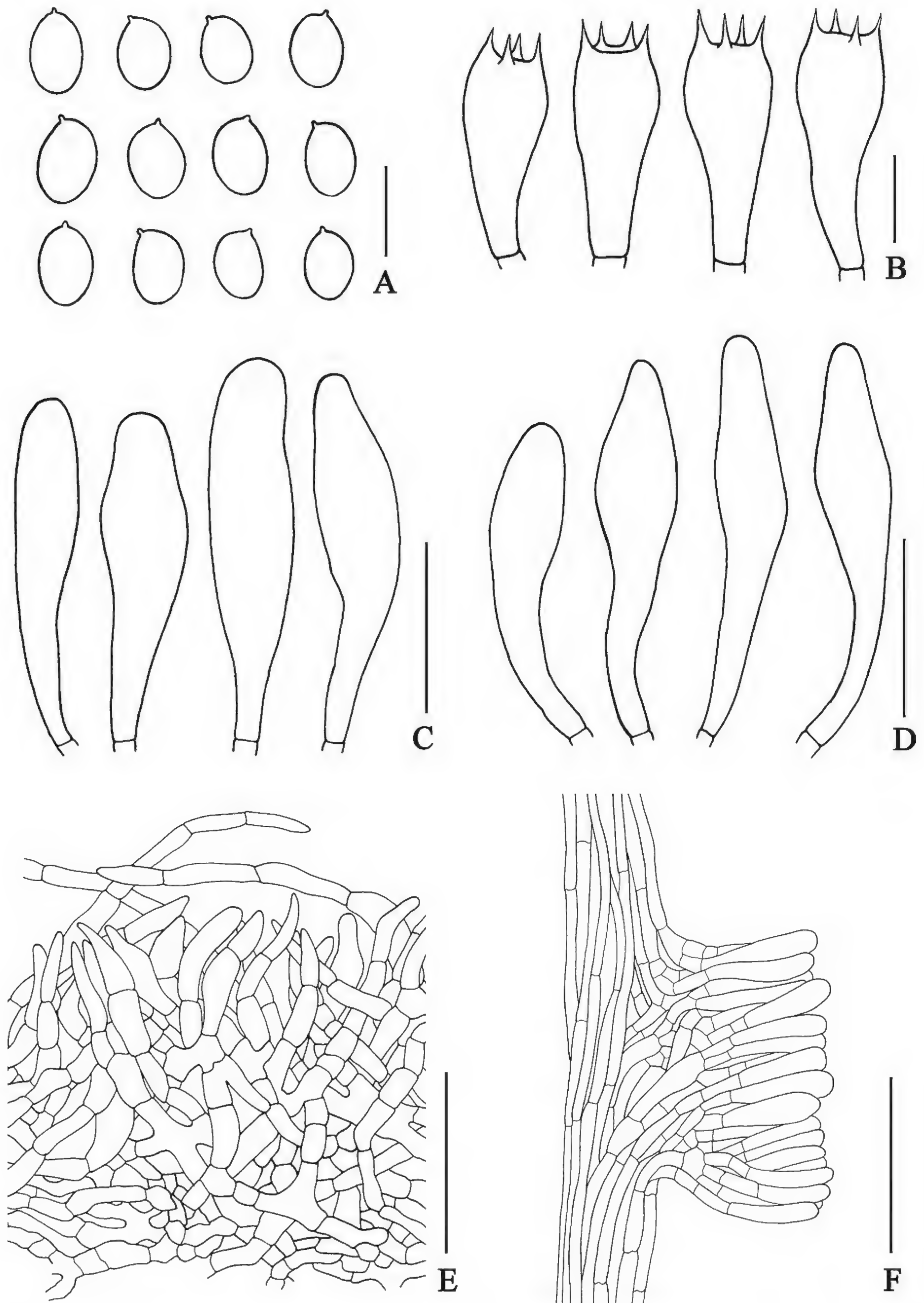


Figure 4. Microscopic features of *Rostrupomyces sisongkhramensis* **A** basidiospores **B** basidia **C** cheilocystidia **D** pleurocystidia **E** pileipellis **F** stipitipellis showing a cluster of narrowly clavate to clavate cells which slightly scattered on the stipe surface. Scale bars: 10 µm (**A–D**); 25 µm (**D–E**); 50 µm (**E–F**). All line drawings were made from SV0155.

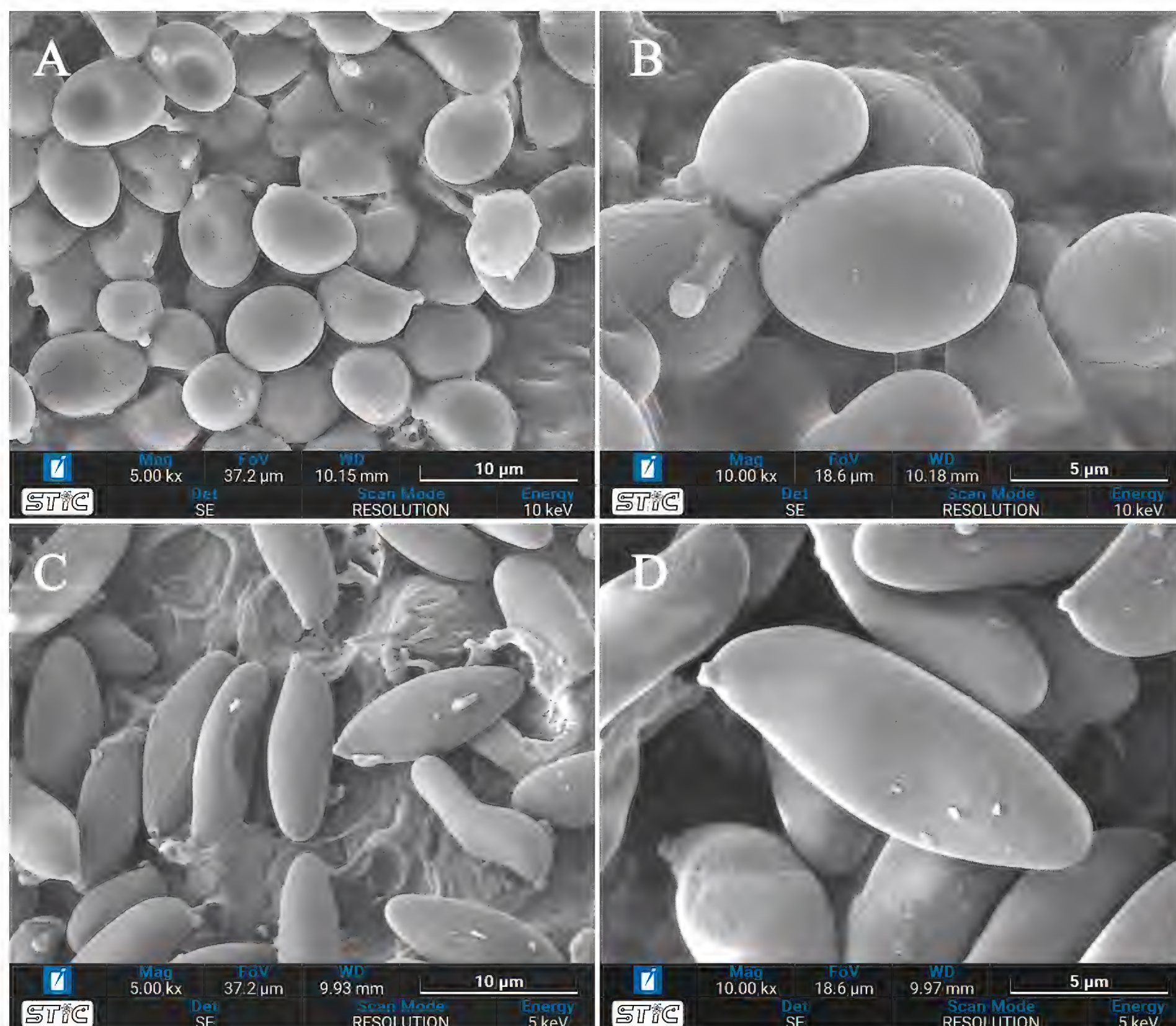


Figure 5. Scanning electron micrographs of basidiospores **A–B** *Rostrupomyces sisongkhramensis* (SV0155) **C–D** *Hemileccinum inferius* (SV0282).

when mature, separable from the pileus context, unchanging when cut. **Pores** 0.2–0.8(1.3) mm wide half-way to the margin, irregularly arranged, roundish then subangular to angular in age; topography subregular, composite pores frequent; color distribution even, when young white (1A1) then yellowish white (4A2) becoming grayish yellow (4B3–5) infrequently with reddish brown spots (7–8E/F8) at places in age, unchanging when touched. **Odour** mild fungoid. **Taste** mild. **Spore print** yellowish brown (5F5) in mass.

Macrochemical reactions: KOH, brownish orange on pileus, yellowish to pale dull orange on pileus context and stipe surface, none or yellowish on stipe context, yellowish brown to brownish orange on hymenium; NH_4OH , yellowish to brownish orange (occasionally with purple aura) on pileus, yellowish to pale orange on stipe surface, yellowish to brownish on hymenium, none or yellowish on pileus context and stipe context.

Spores [591/10/10] (6.3–)6.9–7.9–9.1(–9.8) \times (4.5–)4.8–5.5–6.2(–6.5) μ m $Q = (1.2–)1.29–1.44–1.63(–1.79)$. From the type (6.5–)6.9–7.7–8.8(–9.5) \times (4.7–)5–5.5–6.2(–6.5) μ m, $Q = (1.2–)1.25–1.41–1.54(–1.63)$, $N = 106$, broad-

ly ellipsoid to ellipsoid, thin-walled, smooth under light microscope and SEM, yellowish hyaline in water or KOH, inamyloid. **Basidia** 4-spored, (22–)22–26–31(–31) × (9–)9–11–13(–13) µm, clavate without basal clamp connection, hyaline to yellowish hyaline in KOH; sterigmata up to 4 µm long. **Cheilocystidia** (30–)30–43–58(–59) × (9–)9–11–15(–15) µm, frequent, narrowly fusiform to fusiform with obtuse apex or narrowly utriform, thin-walled, hyaline in KOH. **Pleurocystidia** (33–)33–43–63(–63) × (8–)8–11–13(–13) µm, infrequent, narrowly fusiform to fusiform with obtuse apex, thin-walled, hyaline in KOH. **Hymenophoral trama** subregular to slightly divergent, 38–82 µm wide, with subregular mediostratum 8–24 µm wide, composed of cylindrical, 4–12 µm wide hyphae, hyaline in KOH. **Pileipellis** an intricate trichoderm, 70–130 mm thick, made of moderately interwoven (when young) to loosely interwoven in age, thin-walled, smooth, hyaline hyphae 4–18 mm wide, branching and anastomosing at places; terminal cells 12–65 × 4–18 mm, narrowly fusiform to fusiform to broadly fusiform with slightly acuminate or obtuse apex, hyaline to yellowish pale brown in KOH. **Pileus context** made of strongly interwoven, thin-walled hyphae, up to 12 µm wide, hyaline in KOH. **Stipitipellis** arranged parallel to the surface of the stipe, composed of moderately interwoven, cylindrical, thin-walled, 3–10 µm wide hyphae, anastomosing and branching at places, sparsely scattered with groups of rising cells to clusters (up to 87 µm high) of narrowly clavate to clavate cells (21–36 × 4–9 µm), hyaline to yellowish hyaline in KOH. **Caulocystidia** not seen. **Stipe context** parallelly arranged, composed of moderately interwoven, cylindrical, thin-walled, 3–18 µm wide hyphae, hyaline to yellowish hyaline in KOH. **Clamp connections** not seen in any tissue.

Habitat and distribution. Solitary or in small groups (up to 4 basidiomata), or fasciculate by 2 to 3 basidiomata, on sandy loam to sandy clay loam soil in open dry dipterocarp forest and dipterocarp forest dominated by Dipterocarpaceae trees namely *Anthoshorea roxburghii*, *Dipterocarpus obtusifolius*, *D. tuberculatus*, *D. intricatus*, *Pentacme siamensis*, and *Shorea obtusa* with or without scattered Fagaceae trees. Currently known from the type locality (Nakhon Phanom province), Sisaket and Ubon Ratchathani provinces in northeastern Thailand, and also in Chiang Mai and Chiang Rai provinces in northern Thailand.

Specimens examined. THAILAND, Chiang Mai Province, Muang District, Doi Suthep-Pui National Park, 18°47'39.4"N, 98°55'21.5"E, elev. 915 m, 20 July 2015, *Olivier Raspé*, OR1004 (CMUB, BKF, BR); *ibid.*, 18°48'04.2"N, 98°55'44.3"E, elev. 775 m, 21 July 2015, *Santhiti Vadthanarat*, SV0155 (CMUB, BKF); Mae On District, 18°51'57.4"N, 99°17'22.9"E, elev. 660 m, 1 June 2015, *Olivier Raspé*, OR0915 (CMUB, BR); *ibid.*, 18°51'57.0"N, 99°17'23.0"E, elev. 660 m, 1 June 2015, *Olivier Raspé*, OR0918 (CMUB, BR); *ibid.*, 18°51'57.0"N, 99°17'23.0"E, elev. 660 m, 1 June 2015, *Olivier Raspé*, OR0919 (CMUB, BR); *ibid.*, 18°52'13.0"N, 99°18'25.0"E, elev. 760 m, 15 August 2015, *Santhiti Vadthanarat*, SV0219 (CMUB, BR); *ibid.*, 18°51'57.4"N, 99°17'22.0"E, elev. 700 m, 16 August 2015, *Santhiti Vadthanarat*, SV0225 (CMUB, BR); *ibid.*, 18°51'57.7"N, 99°17'26.5"E, elev. 685 m, 1 June 2017, *Santhiti Vadthanarat*, SV0397 (CMUB, BR); *ibid.*, 18°52'15.6"N, 99°18'11.5"E, elev. 800 m, 11 July 2017, *Olivier Raspé*, OR1392 (CMUB, BR); *ibid.*, 18°52'15.6"N, 99°18'11.5"E, elev. 800 m, 11 July 2017, *Olivier Raspé*, OR1399 (CMUB, BR); *ibid.*, 18°52'16.7"N, 99°18'13.0"E, elev. 800 m, 9 June 2021, *Santhiti Vadthanarat*, SV0512 (CMUB, BR); *ibid.* 18°52'7.9"N, 99°17'42.0"E, elev. 780 m, 10 June 2021, *Santhiti Vadthanarat*, SV0517 (CMUB, BR); *ibid.* 18°52'16.4"N,

99°17'40.5"E, elev. 820 m, 10 June 2021, *Santhiti Vadthanarat*, SV0518 (CMUB, BR); *ibid.* 18°52'12.0"N, 99°17'31.2"E, elev. 700 m, 10 June 2021, *Bhavesh Raghoonundon*, BR0311; *ibid.* 18°52'26.8"N, 99°18'15.5"E, elev. 845 m, 10 June 2021, *Bhavesh Raghoonundon*, BR0313; Chiang Rai Province, Phan District, 19°48'50.0"N, 99°51'57.0"E, elev. 730 m, 22 June 2021, *Bhavesh Raghoonundon*, BR0368; *ibid.* 19°48'50.0"N, 99°51'57.0"E, elev. 730 m, 22 June 2021, *Bhavesh Raghoonundon*, BR0371; Sisaket Province, Kanthararom District, Kok Yang Yai roadside market, 17 September 2016, *Santhiti Vadthanarat*, SV0345 (CMUB); Ubon Ratchathani Province, Trakan Phuet Phon District, Huay Fai, 15°32'44.3"N, 105°10'17.4"E, elev. 165 m, 28 July 2015, *Olivier Raspé*, OR1059 (CMUB, BR).

ITS sequence accession number (SV0155): PP354891.

Notes. The BLAST result based on ITS sequence obtained from one of the examined specimens (voucher SV0155, GenBank accession number PP354891) was 100% identical to the holotype of *X. sisongkhramensis* (voucher BBH 48255, accession number OP462477) which was reported by Tan et al. 2022. This suggested that our collections belonged to *X. sisongkhramensis*. Morphological characters of our collections mostly fit the original description of the species. However, some variations were observed between ours and the original description as follows: Tan et al. (2022) mentioned the absence of cheilocystidia in *X. sisongkhramensis* while we could observe them in our collections; they were narrowly fusiform to fusiform with obtuse apex or narrowly utriform, thin-walled. The protologue mentioned broadly clavate to subclavate (40–60 × 8–15 µm) caulocystidia. However, in our observation only groups of rising terminal cells of shape and size similar to the caulocystidia in Tan et al. (2022), were observed. What Tan et al. (2022) considered as caulocystidia were what we described as undifferentiated terminal cells of the stipitipellis. In the species protologue, the pileipellis and stipitipellis were described as composed of thick-walled hyphae (no measurement mentioned). However, only thin-walled hyphae were observed in our collections.

Rostrupomyces sisongkhramensis is morphologically similar to *Hemileccinum duriusculum* Mei-Xiang Li, Zhu L. Yang & G. Wu, which was recently described from China. The two species share some morphological characters including basidiome size and color, scattering of granular squamules on the stipe surface, pale yellow to grayish yellow hymenophore that is depressed around the stipe apex, and unchanging color in any parts. However, *H. duriusculum* differs by its strikingly venose pileus surface, finer granular squamules on the stipe surface, and subfusiform basidiospores ornamented with irregular warts under SEM (Liu et al. 2024). *Rostrupomyces sisongkhramensis* is also somewhat similar to a European *Leccinum* species originally described from Italy, *Leccinum albstipitatum* den Bakker & Noordel., which has a similar shade of pileus color (light orange), whitish stipe covered with whitish squamules when young to reddish brown in age. However, *L. albstipitatum* can be differentiated by having an inflexed margin which exceeds the hymenophore by up to 4 mm, yellowish white to very pale brown hymenophore that becomes brownish when bruised, a clear blue discoloration of the stipe base when touched, context staining vinaceous then grayish to blackish when cut, smooth fusiform basidiospores, distribution in Europe, and association with *Populus* L. trees (den Bakker and Noordeloos 2005).

Phylogenetically, *R. sisongkhramensis* is closely related to *Rubinosporus auriporus* Vadthanarat, Raspé & Lumyong, the only known species in the genus, which was described from the same region as *Rostrupomyces* (northern Thailand). However, it can be differentiated from *R. sisongkhramensis* by having grayish red to pastel red to reddish brown pileus; even stipe surface with scattered bright yellow to yellowish white to orange to light brown minute squamules; shorter tubes especially when young; golden yellow hymenophore; and the striking dark ruby spore print (Vadthanarat et al. 2022).

***Hemileccinum inferius* Vadthanarat, Raghoonundon & Raspé, sp. nov.**

MycoBank No: 849063

Figs 6, 7

Etymology. “inferius” refers to the only lower part of the stipe ornamented with reticulum

Holotype. THAILAND, Chiang Mai Province, Muang District, Doi Suthep-Pui National Park, 18°47'52.8"N, 98°54'21.2"E, elev. 1,170 m, 1 July 2016, *Santhiti Vadthanarat*, SV0282 (holotype: CMUB, isotype: BKF, MFUB). ITS sequence accession number PP354892.

Diagnosis. *Hemileccinum inferius* can be differentiated from resembling *Hemileccinum* species by a grayish red to reddish brown to dark brown, plane to sub-depressed, subrugulose to pitted pileus; and yellow to yellowish white, cylindrical with subbulbous stipe, with surface even on the upper half and subscabrous to delicately reticulate on the lower half, as well as smooth basidiospores even when observed under SEM.

Description. **Basidiomata** medium-sized. **Pileus** 66–68 mm in diameter, plane to sub-depressed at the centre; **margin** deflexed in age, elastic, slightly exceeding (up to 1 mm); **surface** subrugulose to pitted especially near the margin, dull, moist to slippery when wet, tomentose, grayish red (8B/C3–4) to reddish brown (8D/E4–6) to dark brown to reddish brown (7–8F4–6), unchanging when bruised; **context** 8–10 mm thick half-way to the margin, firm to soft, pale yellow (1A3), slightly brown (7E5) near the pileus surface, light yellow (1A4) above



Figure 6. Fresh basidioma of *Hemileccinum inferius* **A, B** SV0282 (holotype). Scale bars: 1 cm (**A, B**).

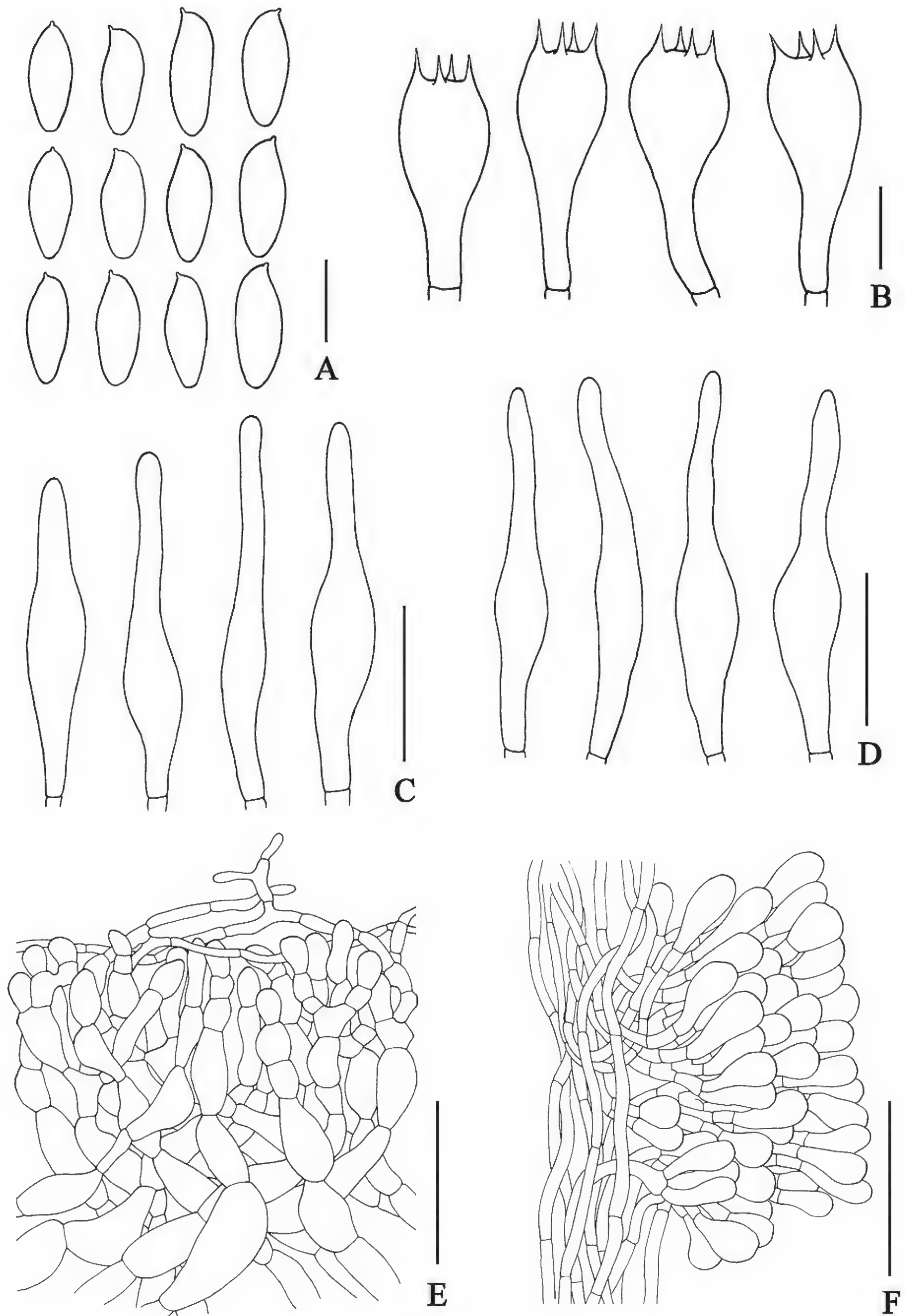


Figure 7. Microscopic features of *Hemileccinum inferius* **A** basidiospores **B** basidia **C** cheilocystidia **D** pleurocystidia **E** pileipellis **F** stipitipellis showing a cluster of clavate to boardy clavate like cells which moderately scattered on the stipitipellis. Scale bars: 10 µm (**A–D**); 25 µm (**D–E**); 50 µm (**E–F**). All drawings were made from holotype type (SV0282).

the hymenium in age, unchanging when cut. **Stipe** 65–76 × 14–18 mm, central, terete, cylindrical with subbulbous base; **surface** even on the upper half then subscabrous to delicately reticulate on the lower half, dull, dry to moist, light yellow (2A4–6) to yellowish white to pale yellow (2A2–3) at the base, occasionally with reddish brown to dark brown spots (8D5–8, 8F7) at places, minutely covered with pale yellow to light brown to dark brown (2A3–4 to 7D/E4, 7F8) squamules on the upper half, slightly fibrillose following a reticulate pattern at the middle of the stipe getting less so to the base, unchanging when bruised; **basal mycelium** white (1A1); **context** solid, firm, pale yellow (2A3–5) especially in the above half near the stipe surface becoming yellowish white (2A2) to off-white at the base, unchanging when cut. **Hymenophore** tubulate, slightly depressed around the stipe, with slightly decurrent tooth, subventricose. **Tubes** 7–8 mm long half-way to the margin, yellow to grayish yellow (2A7 to 2B7) near the pileus context then olive (2E5) near the pores, separable from the pileus context, unchanging when bruised. **Pores** 0.3–0.8(1.2) mm wide at mid-radius, subangular to angular, even, grayish yellow (2B5), unchanging when touched, irregularly arranged; topography subregular. **Odour** mild fungoid. **Taste** mild. **Spore print** olive brown (4E7).

Macrochemical reactions: KOH, brownish orange on pileus and hymenophore, pale orange on pileus context and stipe surface, and stipe context; NH₄OH, brownish orange with purple aura on pileus, yellowish to brownish orange with purple aura on stipe surface, yellowish to greenish or slightly blue on pileus context and stipe context.

Spores [118/2/2] (10.5–)11.4–12.9–14.6(–15.3) × (3.8–)4.2–4.8–5.6(–6.1) µm $Q = (2.06–)2.4–2.68–3.05(–3.32)$. From the type (10.8–)11.5–12.7–14.2(–14.5) × (4.1–)4.3–4.8–5.5(–6.1) µm, $Q = (2.06–)2.33–2.66–3.06(–3.1)$, $N = 68$, narrowly ellipsoid to subcylindrical with a slight suprahilar depression, thin-walled, smooth under light microscope and SEM (Fig. 5C–D), yellowish to brownish hyaline in water, yellowish hyaline in KOH, inamyloid. **Basidia** 4-spored, (23–)24–27–31(–32) × (11–)11–12–14(–14) µm, clavate without basal clamp connection, hyaline to yellowish hyaline in KOH; sterigmata up to 4 µm long. **Cheilocystidia** (30–)34–54–72(–72) × (7–)8–10–14(–14) µm, narrowly fusiform with elongated obtuse apex, frequent, thin-walled, hyaline to yellowish hyaline in KOH. **Pleurocystidia** (34–)34–51–69(–70) × (10–)10–11–13(–13) µm, frequent near the pores, narrowly fusiform with elongated obtuse apex, thin-walled, hyaline to yellowish hyaline in KOH. **Hymenophoral trama** slightly divergent, 62–150 µm wide composed of cylindrical, 4–12 µm wide hyphae, with subregular mediostratum 30–100 µm wide, hyaline in KOH. **Pileipellis** a hyphoepithelium, 80–112 µm thick, the pileipellis composed of ellipsoid to broadly ellipsoid or cylindrical, thin-walled, more or less vertically arranged, occasionally branching or anastomosing, with metablematic, elongated-cylindrical hyphae (2–4 µm wide hyphae), branching or anastomosing at places, hyaline to yellowish hyaline in KOH; terminal cells of 2 types: 1) ellipsoid to broadly ellipsoid, 8–15 × 12–20 µm; and 2) clavate to broadly clavate with obtuse apex, 10–20 × 4–7 µm. **Pileus context** made of moderately interwoven, ellipsoid to broadly ellipsoid, thin-walled hyphae, 10–23 µm wide, hyaline in KOH. **Stipitipellis** arranged parallel to the surface of the stipe (40–50 µm thick), composed of moderately interwoven, cylindrical, thin-walled, 2.5–4 µm wide hyphae, anastomosing at places, moderately scattered with groups of rising cells to clus-

ters (50–60 µm high) of thin-walled clavate to broadly clavate cells (20–30 × 10–15 µm), hyaline to yellowish hyaline in KOH. **Caulocystidia** not seen. **Stipe context** composed of parallel, 8–22 µm wide hyphae, hyaline in KOH. **Clamp connections** not seen in any tissue.

Habitat and distribution. Solitary, on loamy soil in hill evergreen forest dominated by Fagaceae scattered with a few *Dipterocarpus obtusifolius*, at 985–1,170 m elevation. Currently known from Chiang Mai Province, northern Thailand.

Additional specimens examined. THAILAND, Chiang Mai Province, Mae Taeng District, 19°06'59"N, 98°44'23"E, elev. 985 m, 6 June 2021, Bhavesh Raghonundon, BR0260 (MFLU).

Notes. *Hemileccinum inferius* is described based on collections from Thailand. The comparison of the new species with the seven known Asian species follows. *Hemileccinum albidum* differs from *H. inferius* by gray-brown to chrome yellow to ochraceous or golden brown pileus; longer and slender stipe (up to 160 mm); shorter basidiospores (10–12.5 × 4.0–5.5 µm); occurrence at higher elevations (1,968–2,490 m; Li et al. 2021). *Hemileccinum brevisporum* is similar in pileus color but has shorter basidiospores (9–11 × 4–5 µm); and it occurs under Fagaceae and Pinaceae, at higher elevations (1,700–2,120 m; Li et al. 2021). *Hemileccinum duriusculum* is macromorphologically quite similar, but differs by a strongly venose pileus surface, even when young, the absence of reticulum on the lower half of the stipe, as well as shorter cheilo- and pleurocystidia (Liu et al. 2024). *Hemileccinum ferrugineipes* has similar pileus surface and color but can be differentiated by the apparent pale red-brown color on the lower part of the stipe; and shorter basidiospores (11.0–12.5 × 4–5 µm; Li et al. 2021). *Hemileccinum indecorum* is clearly different in having dark red to reddish brown basidiomata with mucilaginous surface densely covered with whitish to dirty white, small conical to subconical to irregularly shaped squamules; incurved margin; and yellowish hymenophore that slowly turns brownish to reddish brown when bruised (Horak 2011; Zeng et al. 2012). *Hemileccinum parvum* has smaller basidiomata (pileus 3.3–3.6 cm diam, stipe 60–97 × 4–9 mm); paler pileus (brownish to yellowish); pale yellow context that slowly turns pale blue when cut (Li et al. 2021). *Hemileccinum rugosum* has paler pileus (light orange to reddish orange); very distinctly rugose to wrinkled pileus surface; and shorter basidiospores (9–13 × 4–5 µm; Wu et al. 2016).

Hemileccinum inferius is also similar to an American species, *H. floridanum*, which has reddish brown to chestnut brown wrinkled and uneven pileus, whitish to pale yellow stipe, white basal mycelium, yellow hymenophore, and smooth basidiospore under both light microscope and SEM. However, the latter species is different by white context that slowly turns yellow often from the margin toward the center, longer basidiospores (10–17 × 4.5–6 µm), likely forms association with oak in northern America (Farid et al. 2021).

Phylogenetically, *H. inferius* was most closely related to *H. hortonii*, *H. rugosum*, and an undescribed specimen (voucher HKAS 53421) from China. *Hemileccinum hortonii*, an American species, can easily be distinguished by its conspicuously pitted pileus, smooth to lightly pruinose stipe that sometimes has delicate reticulation on the upper half, pores that occasionally turn blue on when touched, and slightly longer and narrower basidiospores (12–15 × 3.5–4.5; Kuo and Ortiz-Santana 2020; Farid et al. 2021). For morphological comparison with *H. rugosum* see the above paragraph.

Discussion

In this study, the morphological and phylogenetic evidence highly supported establishing *Rostrupomyces* as a new genus of Boletaceae to accommodate *Xerocomus sisongkhramensis*. The most important morphological characters used to differentiate the new genus from other Boletaceae genera are: subscabrous stipe surface with scattered granulose squamules; hymenophore that is white in young basidiomes and becomes yellow in age; yellowish brown spore print; and broadly ellipsoid to ellipsoid basidiospores with smooth surface.

The character of subscabrous to scabrous stipe surface dotted with scattered granulose squamules is also present in other Boletaceae genera such as *Hemileccinum* (see in notes under *Rostrupomyces*), *Leccinum* Gray, *Leccinellum* Bresinsky & Manfr. Binde, *Rugiboletus* G. Wu & Zhu L. Yang, and *Sutorius* Halling, Nuhn & N.A. Fechner. *Leccinum* can be separated from *Rostrupomyces* by having a white to pallid to light brown hymenophore while *Leccinellum* has yellow hymenophore similar to *Rostrupomyces*. However, both genera are different from *Rostrupomyces* by a more or less pronounced color change of hymenophore, stipe surface, and/or context, which can stain red, brown, yellow, or occasionally blue when bruised. *Leccinellum* and *Leccinum* produce boletoid basidiospores which are also different from *Rostrupomyces*. Moreover, they are phylogenetically distant and placed in another subfamily, the Leccinoideae (den Bakker and Noordeloos 2005; Wu et al. 2016; Xue et al. 2019; Meng et al. 2021). *Rugiboletus* differs from *Rostrupomyces* by its strongly wrinkled pileus (especially when young), yellow or brown or reddish brown hymenophore that is unchanging or turns bluish when bruised, subfusiform basidiospores, and phylogenetically distant and placed in *Pulveroboletus* group (Wu et al. 2015; Kuo and Ortiz-Santana 2020). *Sutorius* Halling, Nuhn & N.A. Fechner, is different in having chocolate to reddish brown or purplish brown basidiomata, grayish or reddish brown or brownish orange hymenophore, context always with scattered reddish or violet or dark brown encrustations that are visible with the naked eye, reddish brown spore deposit, and narrowly ellipsoid to subcylindrical basidiospores (Halling et al. 2012; Vadthanarat et al. 2021). Like *Rugiboletus*, *Sutorius* is phylogenetically distant from *Rostrupomyces*, belonging to the *Pulveroboletus* group (Vadthanarat et al. 2021).

Xerocomoideae genera other than *Rostrupomyces* also produce smooth basidiospores, including *Amylotrama*, *Aureoboletus*, *Alessiaporus*, *Pulchroboletus*, *Rubinosporus*, and *Veloboletus*. Moreover, while most *Xerocomus* and *Phylloporus* species produce basidiospores with bacillate surface, a few species produce smooth basidiospores (Neves and Halling 2010; Wu et al. 2016; Chuankid et al. 2019). However, only *Amylotrama* and *Rubinosporus* present the same shape of basidiospore as *Rostrupomyces*, whereas the others produce more or less oblong to ellipsoid to fusiform basidiospores (Gelardi et al. 2014; Wu et al. 2016; Farid et al. 2017; Frank et al. 2017; Zhang et al. 2019; Crous et al. 2020; Lebel et al. 2022; Vadthanarat et al. 2022). *Amylotrama* comprises two species from Australia, which are completely different from *Rostrupomyces* by their sequestrate basidiomata (Lebel et al. 2022). *Rubinosporus*, differs by having a strikingly thin hymenophore, especially when young; golden yellow hymenophore; and dark ruby spore print (Vadthanarat et al. 2022). *Aureoboletus* differs by the pileus usually having a viscid surface especially when moist; and golden

yellow hymenophore (Wu et al. 201; Zhang et al. 2019). *Alessioporus* is different by its reticulated stipe occasionally with a granular ring-like zone in the middle or lower half of the stipe, golden yellow hymenophore; blue staining of the stipe surface, hymenophore, and context; and distribution in Mediterranean Italy and subtropical USA (Gelardi et al. 2014; Frank et al. 2017). *Pulchroboletus* differs by the stipe surface with scattered red to reddish brown, occasionally with reticulum or longitudinal striations, and with a pseudo-annulus; golden yellow hymenophore; intense blue staining of the hymenophore and context; and occurrence only in Mediterranean Europe and tropical to subtropical America (Gelardi et al. 2014; Farid et al. 2017). The only *Veloboletus* species, is different by its basidiomata with a distinctive universal veil; blue staining of the pileus, stipe, hymenophore, and context, and distribution in Australia (Crous et al. 2020).

Tan et al. (2022) phylogeny was based on ITS and LSU sequences of only *Xerocomus* spp., and *Phylloporus* as outgroup, which resulted in the clustering of *X. sisongkhramensis* in *Xerocomus*. However, our phylogeny based on multiple protein-coding genes (*atp6*, *cox3*, *tef1*, and *rpb2*) and on a much wider taxon sampling of Boletaceae resolved *X. sisongkhramensis* in subfamily Xerocomoideae indeed, but distant from other *Xerocomus* species. Keeping *X. sisongkhramensis* would have rendered the genus polyphyletic. The erection of the new genus *Rostrupomyces*, which can also be morphologically separated from *Xerocomus*, was therefore necessary.

In the phylogeny, *Rostrupomyces* appeared sister to another monotypic genus, *Rubinosporus* (morphological comparison see in notes under *Rostrupomyces sisongkhramensis*). The two genera can be differentiated mainly by the spore print color, and color of hymenophore, two characters that do not vary between species in the same genus in Boletaceae. The characters have been primarily used to differentiate many genera in Boletaceae e.g., *Sutorius*, *Cacaoporus*, *Hourangia*, *Baorangia* (Halling et al. 2012; Wu et al. 2015; Zhu et al. 2015; Vadthanarat et al. 2019b). Additional morphological characters, including pileus color and stipe surface, could be useful to separate them. However, both genera so far comprise only a single species and the pileus color and stipe surface are found to be variable between species within the same genus. For example, in *Boletus* L. and *Tylopilus* P. Karst. the pileus color is variable from white, yellow, brown, orange, green, gray, and purple, and the stipe surface from even to reticulate to strongly reticulate (e.g., Cui et al. 2015; Wu et al. 2016; Li and Yang 2021). Hence, if more species in either of those two genera are described, the comparison between the two genera might need updating.

Rostrupomyces has been found so far on sandy loam to sandy clay loam soils at elevations lower than 1,000 m (165 to 915 m), in open dry dipterocarp and dipterocarp forest mainly dominated by ectomycorrhizal trees in family Dipterocarpaceae genera *Anthoshorea* (*A. roxburghii*), *Dipterocarpus* (*D. obtusifolius*, *D. tuberculatus*, *D. intricatus*), *Pentacme* (*P. siamensis*), and *Shorea* (*S. obtusa*), with scattered Fagaceae trees. In Thailand the Dipterocarpaceae tree species are mainly distributed in lowland (<800 m) to mid-elevation forests (800–1,200 m) whereas Fagaceae trees are mostly distribute in mid-elevation to highland forests (>1,200 m) (Gardner et al. 2007). During our surveys on the diversity of Boletaceae in Thailand, no *Rostrupomyces* collection was found in the forests above 1,000 m, where no *Anthoshorea*, *Dipterocarpus*, *Pentacme*, or

Shorea trees were observed or mentioned as occurring. This suggests that the distribution of *Rostrupomyces* depends on the distribution of the mentioned tree genera, and they are inferred as the associated tree hosts of *Rostrupomyces*. However, a more detailed study is needed to confirm the specificity of its relationship with ectomycorrhizal hosts.

In this study, some specimens of *Rostrupomyces sisongkhramensis* were collected from community forests and the species was found to be consumed by local people, in Ubon Ratchathani and Sisaket provinces in lower northeastern Thailand. It is found on sale on roadsides and local markets, along with other Boletaceae in genera such as *Baorangia*, *Boletus*, *Boletellus*, *Heimioporus*, *Retiboletus*, *Sutorius*, and *Tylopilus*. The species is called “Hed Phueng Waan” in which the words “Hed Phueng” refer to bolete and “Waan” means sweet. It can also be called “Hed Phueng Kaw” in which the words “Kaw” means rice. The same local names are also applied to the other bolete species that are mostly white and have sweet taste after cooking such as *Boletus* spp. In this region a local name can be used for different mushroom species which present similarly striking morphological characters. Conversely, one mushroom species may have more than one local name. *Rostrupomyces sisongkhramensis* is also found in the northern parts of Thailand in Chiang Rai and Chiang Mai provinces. However, during our survey in the region, the species has never been found being collected or on sale for consumption by the locals. The protologue of this species (collections from upper northeastern Thailand) did not mention the edibility (Tan et al. 2022).

To date, 15 *Hemileccinum* species have been described worldwide, among which eight are originally from Asia (China: *H. albidum*, *H. brevisporum*, *H. duriusculum*, *H. ferrugineipes*, *H. parvum*, *H. rugosum*; Singapore: *H. indecorum*; and Thailand: *H. inferius*), two species from France in Europe (*H. depilatum* and *H. impolitum*), four species from North America (*H. floridanum*, *H. hortonii*, *H. rubropunctum*, and *H. subglabripes*), and a single species, *H. brunneotomentosum*, from Belize in Central America (Šutara 2008; Halling et al. 2015; Wu et al. 2016; Kuo and Ortiz-Santana 2020; Nitson and Frank 2020; Farid et al. 2021; Li et al. 2021; Liu et al. 2024). Three *Hemileccinum* species have been previously reported to occur in Thailand, namely *H. depilatum* (reported as *Boletus depilatus* Redeuilh), *H. impolitum* (reported as *B. impolitus* Fr.), and *H. indecorum* (Chandrasrikul et al. 2011; Vadthanarat et al. 2019b). The first two species were originally described from France and were then reported from Thailand based on morphological identification only. As we know that the distribution of Boletaceae species depends on the distribution of their hosts, the ecology and host specificity are important characters in distinguishing species in Boletaceae (den Bakker et al. 2004; Dentinger et al. 2010; Cui et al. 2015; Loizides et al. 2019; Gelardi 2020). It is therefore doubtful that European species are also present in Southeast Asia where the forests are dominated by different tree species or families. Unfortunately, no specimens associated with the reports of occurrence in Thailand are available for molecular analysis to compare with European specimens. Moreover, molecular analysis of several *Hemileccinum* specimens obtained in our study showed none of them belong to those European species. It is therefore reasonable to assume that the identifications of the Thai collections as *H. depilatum* and *H. impolitum* were not correct. The

other recorded species, *H. indecorum* was originally described from Singapore in Southeast Asia (Horak 2011). Specimens collected from Thailand were identified based on both molecular and morphological evidences (Vadthanarat et al. 2019b). However, the full morphological description of this Thai collection has not yet been published. In the future, more detail on the species and more records of *Hemileccinum* will be reported.

Basidiospores with tiny warts and pinholes (when observed under SEM) are typical of *Hemileccinum*. However, a few *Hemileccinum* species produce basidiospores with smooth surface, including the new species (Kuo and Ortiz-Santana et al. 2020; Farid et al. 2021). This kind of exception is also found in other Xerocomoideae genera i.e., in *Phylloporus* and *Xerocomus*, In the latter two genera, most of the species produce basidiospores with bacillate surfaces, but a few produce smooth basidiospores (Neves and Halling 2010; Wu et. al. 2016; Chuankid et al. 2019).

A total of 39 new taxa (4 new genera and 35 new species), including those introduced in this paper, have been originally described from Thailand (Rostrup 1902; Yang et al. 2006; Desjardin et al. 2009; Choeyklin et al. 2012; Neves et al. 2012; Halling et al. 2014; Raspé et al. 2016; Vadthanarat et al. 2018; Chuankid et al. 2019; Phookamsak et al. 2019; Vadthanarat et al. 2019a, 2019b, 2020; Chuankid et al. 2021; Raghoonundon et al. 2021; Vadthanarat et al. 2021; Tan et al. 2022; Vadthanarat et al. 2022; This study). Our study on the diversity of Boletaceae in Thailand is still ongoing and is needed to uncover more new taxa and new distribution records for Thailand.

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Additional information

Conflict of interest

The authors have declared that no competing interests exist.

Ethical statement

No ethical statement was reported.

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Author contributions

Conceptualization: SV, OR. Data curation: BR, SV. Formal analysis: SV. Funding acquisition: OR. Investigation: SV. Methodology: SV. Project administration: OR. Resources: BR, OR, SV. Software: SV. Supervision: OR. Validation: SL, OR. Visualization: SV. Writing – original draft: SV. Writing – review and editing: BR, OR, SL.

Author ORCIDs

Santhiti Vadthanarat  <https://orcid.org/0000-0002-9035-0375>

Bhavesht Raghoonundon  <https://orcid.org/0000-0001-6671-2404>

Olivier Raspé  <https://orcid.org/0000-0002-8426-2133>

Data availability

All of the data that support the findings of this study are available in the main text.

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